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OM protein - protein search, using sw model

February 10, 2005, 17:39:57; Search time 476 Seconds Run on:

(without alignments)

912.812 Million cell updates/sec

Title: US-09-892-287-1

Perfect score: 2020

Sequence: 1 MDVLSPLSFIKVSHVRMQGI.....FQIPPGSLREDPLGEAQPQI 372

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 6959266 seqs, 1168006243 residues

Total number of hits satisfying chosen parameters: 6959266

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : Pending Patents AA Main:*

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Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

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Result		Query				
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1	2020	100.0	372	23	US-09-892-287-1	Sequence 1, Appli
2	2020	100.0	372	35	US-10-990-328-8797	Sequence 8797, Ap
3	2020	100.0	448	1	PCT-US02-17526-14	Sequence 14, Appl
4	2020	100.0	448	27	US-10-161-510-14	Sequence 14, Appl
5	2020	100.0	448	35	US-10-990-328-8795	Sequence 8795, Ap
6	2020	100.0	448	35	US-10-990-328-8796	Sequence 8796, Ap
7	2020	100.0	449	27	US-10-170-205E-32368	Sequence 32368, A
8	2020	100.0	449	35	US-10-990-328-8794	Sequence 8794, Ap
9	2020	100.0	480	21	US-09-724-676-79765	Sequence 79765, A
10	2020	100.0	480	21	US-09-724-676-79776	Sequence 79776, A
11	2020	100.0	480	21	US-09-724-676-79782	Sequence 79782, A
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13	2020	100.0	480	21	US-09-724-676A-79776	Sequence 79776, A
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16	2012	99.6	478	27	US-10-161-510-11	Sequence 11, Appl
17	2010	99.5	372	1	PCT-US02-17382-158	Sequence 158, App
18	2010	99.5	372	1	PCT-US02-17526-13	Sequence 13, Appl
19	2010	99.5	372	27	US-10-161-510-13	Sequence 13, Appl
20	2010	99.5	448	1	PCT-US02-17526-12	Sequence 12, Appl
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22	1991	98.6	518	21	US-09-724-676-79769	Sequence 79769, A
23	1991	98.6	518	21	US-09-724-676-79770	Sequence 79770, A
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35	1966.5	97.4	565	21	US-09-724-676-79787	Sequence 79787, A
36	1966.5	97.4	565	21	US-09-724-676-79788	Sequence 79788, A
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ALIGNMENTS

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RESULT 1
US-09-892-287-1
; Sequence 1, Application US/09892287
    GENERAL INFORMATION:
         APPLICANT: Hillman, Jennifer L.
                    Lal, Preeti
                    Corley, Neil C.
                    Shah, Purvi
         TITLE OF INVENTION: NEW HUMAN PHOSPHATIDYLINOSITOL
                             (4,5) BISPHOSPHATE 5-PHOSPHATASE
        NUMBER OF SEQUENCES: 5
         CORRESPONDENCE ADDRESS:
              ADDRESSEE: Incyte Pharmaceuticals, Inc.
              STREET: 3174 Porter Drive
              CITY: Palo Alto
              STATE: CA
              COUNTRY: USA
              ZIP: 94304
        COMPUTER READABLE FORM:
              MEDIUM TYPE: Diskette
              COMPUTER: IBM Compatible
              OPERATING SYSTEM: DOS
              SOFTWARE: FastSEQ for Windows Version 2.0
         CURRENT APPLICATION DATA:
              APPLICATION NUMBER: US/09/892,287
              FILING DATE: 26-Jun-2001
         PRIOR APPLICATION DATA:
              APPLICATION NUMBER: 09/258,643
              FILING DATE: <Unknown>
        ATTORNEY/AGENT INFORMATION:
              NAME: Billings, Lucy J.
              REGISTRATION NUMBER: 36,749
              REFERENCE/DOCKET NUMBER: PF-0334 US
         TELECOMMUNICATION INFORMATION:
              TELEPHONE: 415-855-0555
              TELEFAX: 415-845-4166
    INFORMATION FOR SEQ ID NO: 1:
         SEQUENCE CHARACTERISTICS:
              LENGTH: 372 amino acids
              TYPE: amino acid
              STRANDEDNESS: single
              TOPOLOGY: linear
         IMMEDIATE SOURCE:
              LIBRARY: BRSTNOT03
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; Sequence 8797, Application US/10990328
; GENERAL INFORMATION:
  APPLICANT: CARGILL, Michele
  TITLE OF INVENTION: POLYMORPHISMS IN NUCLEIC ACID MOLECULES
  TITLE OF INVENTION: ENCODING HUMAN ENZYME PROTEINS, METHODS OF DETECTION AND
  TITLE OF INVENTION: USES THEREOF
  FILE REFERENCE: CL001495
  CURRENT APPLICATION NUMBER: US/10/990,328
  CURRENT FILING DATE: 2004-11-17
  NUMBER OF SEQ ID NOS: 558824
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   ORGANISM: Homo sapiens
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; Sequence 14, Application PC/TUS0217526
; GENERAL INFORMATION:
  APPLICANT: EXELIXIS, INC.
  TITLE OF INVENTION: PIBS AS MODIFIERS OF THE p53 PATHWAY AND METHODS OF USE
  FILE REFERENCE: EX02-074C-PC
  CURRENT APPLICATION NUMBER: PCT/US02/17526
  CURRENT FILING DATE: 2002-06-03
  PRIOR APPLICATION NUMBER: US 60/296,076
  PRIOR FILING DATE: 2001-06-05
  PRIOR APPLICATION NUMBER: US 60/328,605
  PRIOR FILING DATE: 2001-10-10
  PRIOR APPLICATION NUMBER: US 60/357,253
  PRIOR FILING DATE: 2002-02-15
  NUMBER OF SEQ ID NOS: 14
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; Sequence 14, Application US/10161510
; GENERAL INFORMATION:
  APPLICANT: EXELIXIS, INC.
  TITLE OF INVENTION: PIBS AS MODIFIERS OF THE p53 PATHWAY AND METHODS OF USE
  FILE REFERENCE: EX02-074C
  CURRENT APPLICATION NUMBER: US/10/161,510
  CURRENT FILING DATE: 2002-06-03
  PRIOR APPLICATION NUMBER: US 60/296,076
  PRIOR FILING DATE: 2001-06-05
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  PRIOR FILING DATE: 2001-10-10
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  PRIOR FILING DATE: 2002-02-15
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; GENERAL INFORMATION:
  APPLICANT: CARGILL, Michele
  TITLE OF INVENTION: POLYMORPHISMS IN NUCLEIC ACID MOLECULES
  TITLE OF INVENTION: ENCODING HUMAN ENZYME PROTEINS, METHODS OF DETECTION AND
  TITLE OF INVENTION: USES THEREOF
  FILE REFERENCE: CL001495
  CURRENT APPLICATION NUMBER: US/10/990,328
  CURRENT FILING DATE: 2004-11-17
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 GENERAL INFORMATION:
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  TITLE OF INVENTION: POLYMORPHISMS IN NUCLEIC ACID MOLECULES
  TITLE OF INVENTION: ENCODING HUMAN ENZYME PROTEINS, METHODS OF DETECTION AND
  TITLE OF INVENTION: USES THEREOF
  FILE REFERENCE: CL001495
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       181 DTSEKKRKPAWTDRILWRLKRQPCAGPDTPIPPASHFSLSLRGYSSHMTYGISDHKPVSG 240
Qy
           257 DTSEKKRKPAWTDRILWRLKROPCAGPDTPIPPASHFSLSLRGYSSHMTYGISDHKPVSG 316
Db
       241 TFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVNDY 300
Qу
           317 TFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVNDY 376
Db
       301 VSYAWVGDSKVSCSDNLNOVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFOIPPGSL 360
Qу
           377 VSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFQIPPGSL 436
Db
       361 REDPLGEAQPQI 372
Qy
           Db
       437 REDPLGEAQPQI 448
RESULT 7
US-10-170-205E-32368
; Sequence 32368, Application US/10170205E
; GENERAL INFORMATION:
  APPLICANT: ADAMS, Mark
  TITLE OF INVENTION: DEVICES, SUCH AS ARRAYS, COMPRISED OF HUMAN PROTEINS OR
PROTEIN
  TITLE OF INVENTION: CAPTURE AGENTS, AND USES THEREOF
  FILE REFERENCE: CL001381
  CURRENT APPLICATION NUMBER: US/10/170,205E
  CURRENT FILING DATE:
                   2002-06-13
  NUMBER OF SEQ ID NOS: 40312
  SOFTWARE: PatentIn version 3.2
 SEQ ID NO 32368
   LENGTH: 449
   TYPE: PRT
   ORGANISM: Homo sapiens
US-10-170-205E-32368
                    100.0%; Score 2020; DB 27; Length 449;
 Query Match
                           Pred. No. 8.9e-198;
 Best Local Similarity
                    100.0%;
 Matches 372; Conservative
                       0; Mismatches
                                       0; Indels
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         1 MDVLSPLSFIKVSHVRMQGILLLVFAKYQHLPYIQILSTKSTPTGLFGYWGNKGGVNICL 60
Qу
           78 MDVLSPLSFIKVSHVRMOGILLLVFAKYOHLPYIOILSTKSTPTGLFGYWGNKGGVNICL 137
DΡ
        61 KLYGYYVSIINCHLPPHISNNYQRLEHFDRILEMQNCEGRDIPNILDHDLIIWFGDMNFR 120
Qy
           Db
       138 KLYGYYVSIINCHLPPHISNNYQRLEHFDRILEMQNCEGRDIPNILDHDLIIWFGDMNFR 197
       121 IEDFGLHFVRESIKNRCYGGLWEKDQLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDY 180
Qу
           198 IEDFGLHFVRESIKNRCYGGLWEKDQLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDY 257
Dh
       181 DTSEKKRKPAWTDRILWRLKRQPCAGPDTPIPPASHFSLSLRGYSSHMTYGISDHKPVSG 240
Qу
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258 DTSEKKRKPAWTDRILWRLKRQPCAGPDTPIPPASHFSLSLRGYSSHMTYGISDHKPVSG 317
Db
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Qу
           318 TFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVNDY 377
Db
        301 VSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFQIPPGSL 360
Qу
           378 VSYAWVGDSKVSCSDNLNOVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFOIPPGSL 437
Db
Qy
        361 REDPLGEAQPQI 372
           438 REDPLGEAQPQI 449
Db
RESULT 8
US-10-990-328-8794
; Sequence 8794, Application US/10990328
 GENERAL INFORMATION:
  APPLICANT: CARGILL, Michele
  TITLE OF INVENTION: POLYMORPHISMS IN NUCLEIC ACID MOLECULES
  TITLE OF INVENTION: ENCODING HUMAN ENZYME PROTEINS, METHODS OF DETECTION AND
  TITLE OF INVENTION: USES THEREOF
  FILE REFERENCE: CL001495
  CURRENT APPLICATION NUMBER: US/10/990,328
  CURRENT FILING DATE: 2004-11-17
  NUMBER OF SEQ ID NOS: 558824
  SOFTWARE: FastSEO for Windows Version 4.0
 SEO ID NO 8794
   LENGTH: 449
   TYPE: PRT
   ORGANISM: Homo sapiens
US-10-990-328-8794
 Query Match
                     100.0%; Score 2020; DB 35; Length 449;
 Best Local Similarity
                     100.0%; Pred. No. 8.9e-198;
 Matches 372; Conservative
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                                            Indels
                                                        Gaps
                                                               0:
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Qу
           78 MDVLSPLSFIKVSHVRMQGILLLVFAKYQHLPYIQILSTKSTPTGLFGYWGNKGGVNICL 137
Db
Qу
        61 KLYGYYVSIINCHLPPHISNNYORLEHFDRILEMONCEGRDIPNILDHDLIIWFGDMNFR 120
           138 KLYGYYVSIINCHLPPHISNNYQRLEHFDRILEMQNCEGRDIPNILDHDLIIWFGDMNFR 197
Db
        121 IEDFGLHFVRESIKNRCYGGLWEKDQLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDY 180
Qy
           198 IEDFGLHFVRESIKNRCYGGLWEKDQLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDY 257
Db
        181 DTSEKKRKPAWTDRILWRLKRQPCAGPDTPIPPASHFSLSLRGYSSHMTYGISDHKPVSG 240
Qу
           258 DTSEKKRKPAWTDRILWRLKRQPCAGPDTPIPPASHFSLSLRGYSSHMTYGISDHKPVSG 317
Db
        241 TFDI.ELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVNDY 300
Qy
           318 TFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVNDY 377
Db
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301 VSYAWVGDSKVSCSDNLNOVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFQIPPGSL 360
Qy
           378 VSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFQIPPGSL 437
Db
       361 REDPLGEAQPQI 372
Qу
           438 REDPLGEAOPOI 449
Db
RESULT 9
US-09-724-676-79765
; Sequence 79765, Application US/09724676
 GENERAL INFORMATION:
  APPLICANT: Compugen LTD
  TITLE OF INVENTION: Variants of alternative splicing
  FILE REFERENCE: 129181.4 Compugen
  CURRENT APPLICATION NUMBER: US/09/724,676
  CURRENT FILING DATE: 2000-11-28
  NUMBER OF SEQ ID NOS: 97222
  SOFTWARE: PatentIn version 3.2
 SEQ ID NO 79765
   LENGTH: 480
   TYPE: PRT
   ORGANISM: Homo sapiens
US-09-724-676-79765
                    100.0%; Score 2020; DB 21;
                                            Length 480;
 Query Match
                    100.0%; Pred. No. 9.9e-198;
 Best Local Similarity
 Matches 372; Conservative
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                                           Indels
                                                    0;
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Qу
           109 MDVLSPLSFIKVSHVRMQGILLLVFAKYQHLPYIQILSTKSTPTGLFGYWGNKGGVNICL 168
Db
        61 KLYGYYVSIINCHLPPHISNNYQRLEHFDRILEMQNCEGRDIPNILDHDLIIWFGDMNFR 120
Qy
           169 KLYGYYVSIINCHLPPHISNNYQRLEHFDRILEMQNCEGRDIPNILDHDLIIWFGDMNFR 228
Db
        121 IEDFGLHFVRESIKNRCYGGLWEKDOLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDY 180
Qу
           229 IEDFGLHFVRESIKNRCYGGLWEKDOLSIAKKHDPLLREFOEGRLLFPPTYKFDRNSNDY 288
Db
        181 DTSEKKRKPAWTDRILWRLKRQPCAGPD'TPIPPASHFSLSLRGYSSHMTYGISDHKPVSG 240
Qу
           289 DTSEKKRKPAWTDRILWRLKROPCAGPDTPIPPASHFSLSLRGYSSHMTYGISDHKPVSG 348
Db
        241 TFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVNDY 300
QУ
           349 TFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVNDY 408
Db
        301 VSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFQIPPGSL 360
Qу
           409 VSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFQIPPGSL 468
Db
        361 REDPLGEAQPQI 372
Qу
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RESULT 10
US-09-724-676-79776
; Sequence 79776, Application US/09724676
; GENERAL INFORMATION:
  APPLICANT: Compugen LTD
  TITLE OF INVENTION: Variants of alternative splicing
  FILE REFERENCE: 129181.4 Compugen
  CURRENT APPLICATION NUMBER: US/09/724,676
  CURRENT FILING DATE: 2000-11-28
  NUMBER OF SEQ ID NOS: 97222
  SOFTWARE: PatentIn version 3.2
 SEQ ID NO 79776
   LENGTH: 480
   TYPE: PRT
   ORGANISM: Homo sapiens
US-09-724-676-79776
 Query Match
                     100.0%;
                            Score 2020; DB 21; Length 480;
 Best Local Similarity
                     100.0%; Pred. No. 9.9e-198;
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 Matches 372; Conservative
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                                            Indels
                                                              0;
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                                                        Gaps
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Qу
           109 MDVLSPLSFIKVSHVRMQGILLLVFAKYQHLPYIQILSTKSTPTGLFGYWGNKGGVNICL 168
Db
        61 KLYGYYVSIINCHLPPHISNNYORLEHFDRILEMONCEGRDIPNILDHDLIIWFGDMNFR 120
Qy
           Db
        169 KLYGYYVSIINCHLPPHISNNYQRLEHFDRILEMQNCEGRDIPNILDHDLIIWFGDMNFR 228
        121 IEDFGLHFVRESIKNRCYGGLWEKDQLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDY 180
Qу
           229 IEDFGLHFVRESIKNRCYGGLWEKDQLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDY 288
Db
        181 DTSEKKRKPAWTDRILWRLKRQPCAGPDTPIPPASHFSLSLRGYSSHMTYGISDHKPVSG 240
Qу
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Db
        241 TFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVNDY 300
Qу
           349 TFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVNDY 408
Db
Qу
        301 VSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFQIPPGSL 360
           409 VSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFQIPPGSL 468
Dh
        361 REDPLGEAQPQI 372
Qy
           469 REDPLGEAQPQI 480
Db
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RESULT 11

US-09-724-676-79782

[;] Sequence 79782, Application US/09724676

[;] GENERAL INFORMATION:

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APPLICANT: Compugen LTD
  TITLE OF INVENTION: Variants of alternative splicing
  FILE REFERENCE: 129181.4 Compugen
  CURRENT APPLICATION NUMBER: US/09/724,676
  CURRENT FILING DATE: 2000-11-28
  NUMBER OF SEQ ID NOS: 97222
  SOFTWARE: PatentIn version 3.2
 SEO ID NO 79782
   LENGTH: 480
   TYPE: PRT
   ORGANISM: Homo sapiens
US-09-724-676-79782
 Query Match
                     100.0%; Score 2020; DB 21; Length 480;
                     100.0%; Pred. No. 9.9e-198;
 Best Local Similarity
 Matches 372; Conservative
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           109 MDVLSPLSFIKVSHVRMQGILLLVFAKYQHLPYIQILSTKSTPTGLFGYWGNKGGVNICL 168
Db
         61 KLYGYYVSIINCHLPPHISNNYQRLEHFDRILEMQNCEGRDIPNILDHDLIIWFGDMNFR 120
Qу
            169 KLYGYYVSIINCHLPPHISNNYORLEHFDRILEMONCEGRDIPNILDHDLIIWFGDMNFR 228
Db
        121 IEDFGLHFVRESIKNRCYGGLWEKDQLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDY 180
Qy
            229 IEDFGLHFVRESIKNRCYGGLWEKDOLSIAKKHDPLLREFOEGRLLFPPTYKFDRNSNDY 288
Db
        191 DTSEKKRKPAWTDRILWRLKRQPCAGPDTPIPPASHFSLSLRGYSSHMTYGISDHKPVSG 240
Qy
            289 DTSEKKRKPAWTDRILWRLKRQPCAGPDTPIPPASHFSLSLRGYSSHMTYGISDHKPVSG 348
Db
        241 TFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVNDY 300
QУ
            349 TFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVNDY 408
Db
        301 VSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFQIPPGSL 360
QУ
            409 VSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFQIPPGSL 468
Db
        361 REDPLGEAOPOI 372
Qу
            111111111111
        469 REDPLGEAQPQI 480
Db
RESULT 12
US-09-724-676A-79765
; Sequence 79765, Application US/09724676A
; GENERAL INFORMATION:
  APPLICANT: Compugen LTD
  TITLE OF INVENTION: Variants of alternative splicing
  FILE REFERENCE: 129181.4 Compugen
  CURRENT APPLICATION NUMBER: US/09/724,676A
  CURRENT FILING DATE: 2000-11-28
  NUMBER OF SEO ID NOS: 97222
  SOFTWARE: PatentIn version 3.2
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; SEQ ID NO 79765
   LENGTH: 480
   TYPE: PRT
   ORGANISM: Homo sapiens
US-09-724-676A-79765
 Query Match
                     100.0%; Score 2020; DB 21; Length 480;
 Best Local Similarity
                     100.0%; Pred. No. 9.9e-198;
 Matches 372; Conservative
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                                          0; Indels
                                                     0; Gaps
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Qу
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           169 KLYGYYVSIINCHLPPHISNNYQRLEHFDRILEMQNCEGRDIPNILDHDLIIWFGDMNFR 228
Db
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           Db
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QУ
           289 DTSEKKRKPAWIDRILWRLKROPCAGPDTPIPPASHFSLSLRGYSSHMTYGISDHKPVSG 348
Db
        241 TFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVNDY 300
Qу
           349 TFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVNDY 408
Db
        301 VSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFQIPPGSL 360
Qу
           409 VSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFQIPPGSL 468
D'n
        361 REDPLGEAQPQI 372
Qy
           111111111
        469 REDPLGEAQPQI 480
מת
RESULT 13
US-09-724-676A-79776
; Sequence 79776, Application US/09724676A
; GENERAL INFORMATION:
  APPLICANT: Compugen LTD
  TITLE OF INVENTION: Variants of alternative splicing
  FILE REFERENCE: 129181.4 Compugen
  CURRENT APPLICATION NUMBER: US/09/724,676A
  CURRENT FILING DATE: 2000-11-28
  NUMBER OF SEQ ID NOS: 97222
  SOFTWARE: PatentIn version 3.2
 SEQ ID NO 79776
   LENGTH: 480
   TYPE: PRT
   ORGANISM: Homo sapiens
US-09-724-676A-79776
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100.0%; Score 2020; DB 21; Length 480;

Query Match

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Db
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Db
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        289 DTSEKKRKPAWTDRILWRLKRQPCAGPDTPIPPASHFSLSLRGYSSHMTYGISDHKPVSG 348
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Qу
           Db
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Qу
           Db
        409 VSYAWVGDSKVSCSDNLNOVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPF0IPPGSL 468
        361 REDPLGEAOPOI 372
QУ
           469 REDPLGEAQPQI 480
Dh
RESULT 14
US-09-724-676A-79782
; Sequence 79782, Application US/09724676A
; GENERAL INFORMATION:
  APPLICANT: Compugen LTD
  TITLE OF INVENTION: Variants of alternative splicing
  FILE REFERENCE: 129181.4 Compugen
  CURRENT APPLICATION NUMBER: US/09/724,676A
  CURRENT FILING DATE: 2000-11-28
  NUMBER OF SEQ ID NOS: 97222
  SOFTWARE: PatentIn version 3.2
 SEO ID NO 79782
   LENGTH: 480
   TYPE: PRT
   ORGANISM: Homo sapiens
US-09-724-676A-79782
 Query Match
                     100.0%; Score 2020; DB 21;
                                             Length 480;
 Best Local Similarity
                     100.0%; Pred. No. 9.9e-198;
 Matches 372; Conservative 0; Mismatches
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           Db
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            Db
        289 DTSEKKRKPAWTDRILWRLKROPCAGPDTPIPPASHFSLSLRGYSSHMTYGISDHKPVSG 348
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Qy
            Db
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Qy
            409 VSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFQIPPGSL 468
Db
        361 REDPLGEAQPQI 372
Qу
            469 REDPLGEAQPQI 480
Db
RESULT 15
PCT-US02-17526-11
; Sequence 11, Application PC/TUS0217526
 GENERAL INFORMATION:
  APPLICANT: EXELIXIS, INC.
  TITLE OF INVENTION: PIBS AS MODIFIERS OF THE p53 PATHWAY AND METHODS OF USE
  FILE REFERENCE: EX02-074C-PC
  CURRENT APPLICATION NUMBER: PCT/US02/17526
  CURRENT FILING DATE: 2002-06-03
  PRIOR APPLICATION NUMBER: US 60/296,076
  PRIOR FILING DATE: 2001-06-05
  PRIOR APPLICATION NUMBER: US 60/328,605
  PRIOR FILING DATE: 2001-10-10
  PRIOR APPLICATION NUMBER: US 60/357,253
  PRIOR FILING DATE: 2002-02-15
  NUMBER OF SEO ID NOS: 14
  SOFTWARE: PatentIn version 3.1
; SEO ID NO 11
   LENGTH: 478
   TYPE: PRT
   ORGANISM: Homo sapiens
   FEATURE:
   NAME/KEY: misc feature
   LCCATION: (56)..(56)
   OTHER INFORMATION: "X" is any amino acid
PCT-US02-17526-11
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                            Score 2012; DB 1; Length 478;
 Best Local Similarity
                      99.7%;
                             Pred. No. 6.5e-197;
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Qy

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Db	167	KLYGYYVSIINCHLPPHISNNYQRLEHFDRILEMQNCEGRDIPNILDHDLIIWFGDMNFR	226
Qy	121	<pre>IEDFGLHFVRESIKNRCYGGLWEKDQLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDY</pre>	180
Db	227	IEDFGLHFVRESIKNRCYGGLWEKDQLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDY	286
Qy	181	DTSEKKRKPAWTDRILWRLKRQPCAGPDTPIPPASHFSLSLRGYSSHMTYGISDHKPVSG	240
Db	287	DTSEKKRKPAWTDRILWRLKRQACAGPDTPIPPASHFSLSLRGYSSHMTYGISDHKPVSG	346
Qy	241	TFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVNDY	300
Db	347		406
Qу	301	VSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFQIPPGSL	360
Db	407	VSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFQIPPGSL	466
Qy	361	REDPLGEAQPQI 372	
Db	467	REDPLGEAQPQI 478	

Search completed: February 10, 2005, 18:01:44 Job time : 477 secs

GenCore version 5.1.6
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OM protein - protein search, using sw model

Run on: February 10, 2005, 17:47:17; Search time 23 Seconds (without alignments)

592.366 Million cell updates/sec

Title: US-09-892-287-1

Perfect score: 2020

Sequence: 1 MDVLSPLSFIKVSHVRMQGI......FQIPPGSLREDPLGEAQPQI 372

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 144007 seqs, 36624764 residues

Total number of hits satisfying chosen parameters: 144007

Minimum DB seq length: 0

Maximum DB seq length: 200000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : Pending_Patents_AA_New:*

1: /cgn2_6/ptodata/2/paa/PCT_NEW_COMB.pep:*
2: /cgn2_6/ptodata/2/paa/US06_NEW_COMB.pep:*
3: /cgn2_6/ptodata/2/paa/US07_NEW_COMB.pep:*
4: /cgn2_6/ptodata/2/paa/US08_NEW_COMB.pep:*
5: /cgn2_6/ptodata/2/paa/US09_NEW_COMB.pep:*
6: /cgn2_6/ptodata/2/paa/US10_NEW_COMB.pep:*
7: /cgn2_6/ptodata/2/paa/US11_NEW_COMB.pep:*
8: /cgn2_6/ptodata/2/paa/US60_NEW_COMB.pep:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

		ક			SOFFARIES	
Result		Query				
No.	Score		Length	DB	ID	Description
1		16.2	908	1	PCT-US04-42360-1594	Sequence 1594, Ap
2	104.5	5.2	458	6	US-10-450-763-57306	Sequence 57306, A
3	90.5	4.5	497	6	US-10-450-763-33366	Sequence 33366, A
4	89.5	4.4	873	6	US-10-496-011-6	Sequence 6, Appli
5	89	4.4	1048	6	US-10-450-763-52226	Sequence 52226, A
6	88	4.4	551	6	US-10-450-763-50598	Sequence 50598, A
7	86.5	4.3	815	6	US-10-496-011-67	Sequence 67, Appl
8	36.5	4.3	851	6	US-10-496-011-3	Sequence 3, Appli
9	85	4.2	321	8	US-60-643-717-11816	Sequence 11816, A
10	85	4.2	512	6	US-10-450-763-34667	Sequence 34667, A
11	85	4.2	735	7	US-11-021-951-123	Sequence 123, App
12	85	4.2	1388	6	US-10-450-763-51703	Sequence 51703, A
13	84.5	4.2	716	6	US-10-450-763-43858	Sequence 43858, A
14	84.5	4.2	731	6	US-10-450-763-48706	Sequence 48706, A
15	84	4.2	380	1	PCT-US04-17965-2224	Sequence 2224, Ap
16	83	4.1	565	6	US-10-450-763-39254	Sequence 39254, A
17	83	4.1	738	6	US-10-450-763-48671	Sequence 48671, A
18	83	4.1	746	6	US-10-450-763-41065	Sequence 41065, A
19	83	4.1	746	6	US-10-450-763-52958	Sequence 52958, A
20	83	4.1	851	6	US-10-450-763-56207	Sequence 56207, A
21	83	4.1	1404	6	US-10-450-763-34552	Sequence 34552, A
22	82.5	4.1	515	6	US-10-450-763-31949	Sequence 31949, A
. 23	82.5	4.1	1114	1	PCT-US04-42360-604	Sequence 604, App
24	82	4.1	376	6	US-10-450-763-39711	Sequence 39711, A
25	82	4.1	940	6	US-10-450-763-32557	Sequence 32557, A
26	81.5	4.0	243	7	US-11-031-175-15153	Sequence 15153, A
27	81	4.0	500	6	US-10-450-763-58159	Sequence 58159, A
23	81	4.0	501	6	US-10-450-763-59146	Sequence 59146, A
29	81	4.0	2114	6	US-10-450-763-39435	Sequence 39435, A
30	81	4.0	2356	6	US-10-450-763-39431	Sequence 39431, A
31	80.5	4.0	535	8	US-60-643-717-1969	Sequence 1969, Ap
32	80.5	4.0	625	7	US-11-031-175-12183	Sequence 12183, A
33	80.5	4.0	631	6	US-10-450-763-52920	Sequence 52920, A
34	80.5	4.0	796	7	US-11-033-545-386	Sequence 386, App
35	80	4.0	633	6	US-10-450-763-57583	Sequence 57583, A
36	80	4.0	1065	6	US-10-450-763-56125	Sequence 56125, A

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80
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37
                                                   Sequence 58750, A
     79.5
                                                   Sequence 3617, Ap
38
            3.9 264 7 US-11-027-399-3617
     79.5
                  264 7 US-11-027-843-3617
                                                   Sequence 3617, Ap
39
            3.9
                  264 7
                                                   Sequence 3617, Ap
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40
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                          US-11-027-878-3617
     79.5
            3.9
                 264 7
                                                   Sequence 3617, Ap
41
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                                                   Sequence 3617, Ap
42
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                                                   Sequence 3617, Ap
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     79.5
           3.9 264 7 US-11-027-879-3617
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                                                   Sequence 3617, Ap
45
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ALIGNMENTS

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RESULT 1
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; Sequence 1594, Application PC/TUS0442360
; GENERAL INFORMATION:
  APPLICANT: The General Hospital Corporation doing business as Massachusetts
General
  APPLICANT: Hospital / Bayer AG
  TITLE OF INVENTION: Nucleotide sequences involved in pain
; FILE REFERENCE: 17633/2048
  CURRENT APPLICATION NUMBER: PCT/US04/42360
  CURRENT FILING DATE: 2004-12-17
; PRICE APPLICATION NUMBER: 60/531,341
; PRIOR FILING DATE: 19-Dec-2003
; NUMBER OF SEQ ID NOS: 2587
 SOFTWARE: Perl script
; SEQ ID NO 1594
   LENGTH: 908
   TYPE: PRT
; ORGANISM: Rattus norvegicus
  PUBLICATION INFORMATION:
   DATABASE ACCESSION NUMBER: Refseq / NP 445928
   DATABASE ENTRY DATE: 2003-10-05
PCT-US04-42360-1594
                       16.2%; Score 328; DB 1; Length 908;
 Query Match
 Best Local Similarity
                       33.6%; Pred. No. 1.5e-21;
 Matches 84; Conservative 44; Mismatches 106; Indels 16; Gaps
                                                                     6;
          9 FIKVSHVRMQGILLLVFAKYQHLPYIQILSTKSTPTGLFGYWGNKGGVNICLKLYGYYVS 68
Qу
            :: :: :: |: | | | | : | | | |:|: :: : | | : | | | | | | | | | | | : : : :
        225 YVLLASEQLVGVCLFVFIRPQHAPFIRDVAVDTVKTGMGGATGNKGAVAIRMLFHTTSLC 284
Db
         69 IINCHLPPHISNNYQRLEHFDRIL-EMONCEGRDIFNILDHDLIIWFGDMNFRIEDFGLH 127
Qу
                   Db
        285 FVCSHFAAGOSOVKERNEDFVEIARKLSFPMGR---MLFSHDYVFWCGDFNYRI-DLPNE 340
        128 FVRESIKNRCYGGLWEKDQLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDYDTSEKKR 187
Qу
             341 EVKELIRQQNWDSLIAGDQLINQKNAGQIFRGFLEGKVTFAPTYKYDLFSEDYDTSEKCR 400
Db
         188 KPAWTDRILWRLKRQPC--AGPDTPIPPASHFSLSLRGYS----SHMTYG-----ISDHK 236
Qу
             Db
         401 TPAWTDRVLWRRRKWPFDRSAEDLDLLNASFQDESKILYTWTPGTLLHYGRAELKTSDHR 460
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237 PVSGTFDLEL 246
Qу
         | | | | :::
        461 PVVALIDIDI 470
Db
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US-10-450-763-57306
; Sequence 57306, Application US/10450763
; GENERAL INFORMATION:
; APPLICANT: Hyseq, Inc
; TITLE OF INVENTION: NOVEL NUCLEIC ACIDS AND POLYPEPTIDES
; FILE REFERENCE: 790CIP3/US
; CURRENT APPLICATION NUMBER: US/10/450,763
  CURRENT FILING DATE: 2003-06-11
  PRIOR APPLICATION NUMBER: PCT/US01/08631
  PRIOR FILING DATE: 2001-03-30
; PRIOR APPLICATION NUMBER: 09/540,217
  PRIOR FILING DATE: 2000-03-31
  PRIOR APPLICATION NUMBER: 09/649,167
 PRIOR FILING DATE: 2000-08-23
; NUMBER OF SEQ ID NOS: 60736
  SOFTWARE: Custom
; SEQ ID NO 57306
; LENGTH: 458
   TYPE: PRT
  ORGANISM: Homo sapiens
   FEATURE:
   NAME/KEY: DOMAIN
   LOCATION: (235)..(278)
   OTHER INFORMATION: 1 kw MIND INHIBITOR DIVISION CONTROL domain identified by
   OTHER INFORMATION: eMATRIX, accession number DM01756B, p-value=4.822e-15,
raw score of
  OTHER INFORMATION: 22.91
   FEATURE:
   NAME/KEY: DOMAIN
   LOCATION: (200)..(305)
   OTHER INFORMATION: ParA family ATPase domain identified by PFam, accession
name
; OTHER INFORMATION: ParA, E-value=4.9e-24, PFam score of 93.3
US-10-450-763-57306
 Query Match
                      5.2%; Score 104.5; DB 6; Length 458;
 Best Local Similarity 20.9%; Pred. No. 0.12;
 Matches 71; Conservative 51; Mismatches 133; Indels 85; Gaps 17;
         46 LFGYWGNKGGVNICLKLYGYYVSIINCHLPPHISNNYQRLEHFDRILEMQNCEGRDIPNI 105
Qу
            108 VIGVAAHKGGV-----YKTSVSVHLAQDLAL-----KGLRVLLVEGNDPQGT 149
        106 --LDHDLIIWFGDMNFRIEDFGLHF-----VRESIKNRCYGGL-WEKDQLSIAKKHDP 155
QУ
              150 ASMYHG---WVPDLHIHAEDTLLPFYLGEKDDVTYAIKPTCWPGLDIIPSCLALHRIETE 206
D'n
        156 LLREFQEGRLLFPP----TYKFDRNSNDYDTSEKKRKP---AWTDRILWRLKRQPCAGPD 208
Qу
            207 LMGKFDEGKLPTDPHLMLRLAIETVAHDYDVIVIDSAPNLGIGTINVV------CAADV 259
Db
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Qу
        209 TPIP-PASHFSLSLRGYSSHMTYGISDHKPVSGTFDLELKPLVSAPLIVLMPEDLWTVEN 267
             260 LIVPTPAELFD-----YTSALQF------FDMLRDLLKNVDLKGFEPDKKWLAGH 303
Db
        268 DMMVSYSSTSDFPSSPWDWIGL-YKVGLRDVNDYVSYAWV----GDSKVSCSDNLNQVYI 322
Qу
            304 EQYIFYAN-SELPKNPRGGLGVRYPGGPHNVPDNLD--WVHLLYNGGQAPCMKSNNALIV 360
Db
        323 DISNIPTTEDEFLLCYYSNSLRSVVGISRPFQIPPGSLRE 362
Qу
                                : || ||:
Db
        361 ILGTVTL------DAVGIGLVMPVLPGLLRD 385
RESULT 3
US-10-450-763-33366
; Sequence 33366, Application US/10450763
; GENERAL INFORMATION:
; APPLICANT: Hyseq, Inc
; TITLE OF INVENTION: NOVEL NUCLEIC ACIDS AND POLYPEPTIDES
; FILE REFERENCE: 790CIP3/US
  CURRENT APPLICATION NUMBER: US/10/450,763
  CURRENT FILING DATE: 2003-06-11
 PRIOR APPLICATION NUMBER: PCT/US01/08631
  PRIOR FILING DATE: 2001-03-30
; PRIOR APPLICATION NUMBER: 09/540,217
; PRIOR FILING DATE: 2000-03-31
; PRIOR APPLICATION NUMBER: 09/649,167
; PRIOR FILING DATE: 2000-08-23
; NUMBER OF SEQ ID NOS: 60736
; SOFTWARE: Custom
; SEQ ID NO 33366
  LENGTH: 497
  TYPE: PRT
  ORGANISM: Homo sapiens
  FEATURE:
  NAME/KEY: DOMAIN
  LOCATION: (235)..(278)
   OTHER INFORMATION: 1 kw MIND INHIBITOR DIVISION CONTROL domain identified by
   OTHER INFORMATION: eMATRIX, accession number DM01756B, p-value=4.822e-15,
raw score of
  OTHER INFORMATION: 22.91
   FEATURE:
   NAME/KEY: DOMAIN
   LOCATION: (200)..(307)
   OTHER INFORMATION: ParA family ATPase domain identified by PFam, accession
;
name
; OTHER INFORMATION: ParA, E-value=6.5e-23, PFam score of 89.5
US-10-450-763-33366
                       4.5%; Score 90.5; DB 6; Length 497;
 Query Match
 Best Local Similarity 20.5%; Pred. No. 2.4;
 Matches 84; Conservative 59; Mismatches 138; Indels 129; Gaps
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Qу
            108 VIGVAAHKGGV-----YKTSVSVHLAQDLAL-----KGLRVLLVEGNDPQGT 149
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Qу
            : | | | :: | | | :| | :: :
        150 ASMYHG---WVPDLHIHAEDTLLPFYLGEKDDVTYAIKPTCWPGLDIIPSCLALHRIETE 206
Db
        156 LLREFQEGRLLFPP----TYKFDRNSNDYDTSEKKRKP---AWTDRILWRLKRQPCAGPD 208
Qу
            ]::|||:|| :: ||
        207 LMGKFDEGKLPTDPHLMLRLAIETVAHDYDVIVIDSAPNLGIGTINVV------CAADV 259
Db
        209 TPIP-PASHF------SLSLRGYSSHMTYGISDHKPVSGTFDLELKPLV 250
QУ
             : | | | |
                                   :: |:|: : :: |:| |
        260 LIVPTPAELFDYTSALOFFDMLRDLLKNVDLKGFEPDTSVSTVTLPSLTLTLNL-LPPSY 318
Db
        251 SAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSP----- 283
Qу
             319 KDPCDYIGP--TWTIODNLPISRSLT--LTTSPKFFLPSERLSSSWLDVRHIEKYVDQGK 374
Db
        284 -----WDW-----IG----LYKVGLRD----VNDYVSYAWVGDSKVSCSDNLNQV- 320
QУ
                 375 SGTRELLWSWAQKNKTIGDLLQVLQEMGHRRAIHLITNYVRHA-VTKSATATATGYQEVL 433
Db
Qу
        321 ----YIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFQIPPGSLRED-PL 365
               434 NSEQILDLTKAP---EVFAIPIQGNRLRGVVLVVGRFGV--SQLKEEHPL 478
Db
RESULT 4
US-10-496-011-6
; Sequence 6, Application US/10496011
; GENERAL INFORMATION:
; APPLICANT: BOURGERON, THOMAS
; APPLICANT: JAMAIN, STEPHANE
; APPLICANT: QUACH, HELENE
; APPLICANT: BETANCUR, CATALINA
; APPLICANT: LEBOYER, MARION
 APPLICANT: GILLBERG, CHRISTOPHER
  TITLE OF INVENTION: POLYNUCLEOTIDE AND PROTEIN INVOLVED IN SYNAPTOGENESIS,
VARIANTS
; TITLE OF INVENTION: THEREOF, AND THEIR THERAPEUTIC AND DIAGNOSTIC USES
; FILE REFERENCE: 253820US0XPCT
; CURRENT APPLICATION NUMBER: US/10/496,011
; CURRENT FILING DATE: 2004-05-28
; PRIOR APPLICATION NUMBER: PCT/FR02/04134
; PRIOR FILING DATE: 2002-11-28
 PRIOR APPLICATION NUMBER: CA2364106
  PRIOR FILING DATE: 2001-11-30
; NUMBER OF SEQ ID NOS: 98
 SOFTWARE: PatentIn version 3.3
; SEQ ID NO 6
  LENGTH: 873
   TYPE: PRT
  ORGANISM: Homo sapiens
US-10-496-011-6
                      4.4%; Score 89.5; DB 6; Length 873;
 Query Match
 Best Local Similarity 18.8%; Pred. No. 6.4;
 Matches 85; Conservative 59; Mismatches 134; Indels 173; Gaps
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106 --LDHDLIIWFGDMNFRIEDFGLHF------VRESIKNRCYGGL-WEKDQLSIAKKHDP 155

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5 SPLSFIKVSH-----VRMOGILLLVFA-KYOHLPYIOILSTKSTPTGLFGYWGNKG 54
QУ
           Db
        315 SCVSLLTLSHYSEGLFQKAIIQSGTALSSWAVNYQPAKYTRILA------DKV 361
        55 GVNI-----PHISNNYORLEH 87
Qу
                   362 GCNMLDTTDMVECLKNKNYKELIQQTITPATYHIAFGPVIDGDVIPDDPQILMEQGEFLN 421
Db
Qу
        88 FDRILEMONCEG-RDIPNILDHDLIIWFGDMNFRIEDF-----GLHFVRESIKNRC 137
           :|:|: ||: ||:|:|
        422 YDIMLGVNQGEGLKFVDGIVDNEDGVTPNDFDFSVSNFVDNLYGYPEGKDTLRETIK-FM 480
Db
        138 YGGLWEKDQLSIAKKHDPLLREFQEGRLLFP-----PTYKFDRNSNDYDTSE 184
Qу
           481 YTDWADKENPETRRK--TLVALFTDHQWVAPAVATADLHAQYGSPTYFY----AFYHHCQ 534
Dh
QУ
        185 KKRKPAWTDRILWRLKRQPCAGPDTPIPPASHFSLSLRGYSSHMTYGISDHKPVSGTFDL 244
            : ||:| |
                                     | | :||
        535 SEMKPSWAD------ 556
Db
Qу
        245 ELKPLVSAPLIVLMPEDLWTV---ENDMMVS-----YSSTSDFPSSPWDWIGLYKV 292
             Db
        557 ---PMIG-----PTELFSCNFSKNDVMLSAVVMTYWTNFAKTGD-PNOPVP----- 598
        293 GLRDV------NDYVSYAWVGDSKVSCSDNLNOVYIDISNIPTTEDEF------L 335
Qу
            599 --QDTKFIHTKPNRFEEVAW---SKYNPKD---QLYLHIGLKPRVRDHYRATKVAFWLEL 650
Db
        336 LCYYSNSLRSVVGISRPFQIPPGSLREDPLG 366
Qу
           :: | :: | : | |
Db
        651 VPHLHNLNEIFQYVSTTTKVPPPDMTSFPYG 681
RESULT 5
US-10-450-763-52226
; Sequence 52226, Application US/10450763
; GENERAL INFORMATION:
  APPLICANT: Hyseq, Inc
  TITLE OF INVENTION: NOVEL NUCLEIC ACIDS AND POLYPEPTIDES
  FILE REFERENCE: 790CIP3/US
  CURRENT APPLICATION NUMBER: US/10/450,763
  CURRENT FILING DATE: 2003-06-11
  PRIOR APPLICATION NUMBER: PCT/US01/08631
  PRIOR FILING DATE: 2001-03-30
  PRIOR APPLICATION NUMBER: 09/540,217
  PRIOR FILING DATE: 2000-03-31
  PRIOR APPLICATION NUMBER: 09/649,167
  PRIOR FILING DATE: 2000-08-23
  NUMBER OF SEQ ID NOS: 60736
  SOFTWARE: Custom
; SEQ ID NO 52226
   LENGTH: 1048
   TYPE: PRT
   ORGANISM: Homo sapiens
   FEATURE:
   NAME/KEY: DOMAIN
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LOCATION: (410)..(439)
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    OTHER INFORMATION: eMATRIX, accession number DM00653A, p-value=7.207e-27,
 raw score of
    OTHER INFORMATION: 20.40
    FEATURE:
    NAME/KEY: DOMAIN
    LOCATION: (651)..(765)
    OTHER INFORMATION: ParA family ATPase domain identified by PFam, accession
 name
 ; OTHER INFORMATION: ParA, E-value=2.7e-28, PFam score of 107.5
 US-10-450-763-52226
  Query Match
                        4.4%; Score 89; DB 6; Length 1048;
  Best Local Similarity 21.4%; Pred. No. 9;
  Matches 75; Conservative 47; Mismatches 112; Indels 116; Gaps
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 Qу
             553 VIGVAAHKGGV-----YKTSVSVHLAQDLAL-----KGLRVLLVEGNDPQGT 599
Db
         106 --LDHDLIIWFGDMNFRIEDFGLHF-----VRESIKNRCYGGL-WEKDQLSIAKKHDP 155
 Qу
              600 ASMYHG---WVPDLHIHAEDTLLPFYLGEKDDVTYAIKPTCWPGLDIIPSCLALHRIETE 656
 Db
         156 LLREFQEGRLLFPP----TYKFDRNSNDYDTSEKKRKP---AWTDRILWRLKRQPCAGPD 208
 Qу
             1::|||:| :::||
         657 LMGKFDEGKLPTDPHLMLRLAIETVAHDYDVIVIDSAPNLGIGTINVV------CAADV 709
 Db
         209 TPIP-PASHFSLSLRGYSSHMTYGISDHKPVSGTFDLELKPLVSAPLIVLMPEDLWTVEN 267
 Qу
                                          :| || | :| : :
         710 LIVPTPAELFD-----YTSALOF------FDM-LRDL-----LKNVDLKGFEP 745
 Db
         268 D--MMVSYSSTSDFPSSPW----- 293
 Qу
             | :::: | |: |||
                                               |\cdot|\cdot|
         746 DVRILLTKYSNSNGSQSPWMEEQIRDAWGSMVLKNVVRETDEVG--KVAEYNGYHFVFAL 803
 Db
         294 -----IRDVNDYVSYAWVGDSKVSCSDNLNQVYID----ISNIPTTEDE 333
 Qу
                    804 AGSPKDAVDTSIYMFYQKVGDNSIDSWKNAGRVFKDSDKFVANDPILKDQ 853
 Db
 RESULT 6
 US-10-450-763-50598
 ; Sequence 50598, Application US/10450763
 ; GENERAL INFORMATION:
 ; APPLICANT: Hyseq, Inc
  TITLE OF INVENTION: NOVEL NUCLEIC ACIDS AND POLYPEPTIDES
  FILE REFERENCE: 790CIP3/US
   CURRENT APPLICATION NUMBER: US/10/450,763
   CURRENT FILING DATE: 2003-06-11
   PRIOR APPLICATION NUMBER: PCT/US01/08631
   PRIOR FILING DATE: 2001-03-30
 ; PRIOR APPLICATION NUMBER: 09/540,217
  PRIOR FILING DATE: 2000-03-31
  PRIOR APPLICATION NUMBER: 09/649,167
   PRIOR FILING DATE: 2000-08-23
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NUMBER OF SEQ ID NOS: 60736
 SOFTWARE: Custom
; SEQ ID NO 50598
   LENGTH: 551
   TYPE: PRT
   ORGANISM: Homo sapiens
   FEATURE:
   NAME/KEY: misc feature
   LOCATION: (1)...(551)
   OTHER INFORMATION: Xaa = X or * as defined in Table 2
US-10-450-763-50598
  Query Match
                       4.4%; Score 88; DB 6; Length 551;
  Best Local Similarity 22.8%; Pred. No. 4.7;
  Matches 64; Conservative 39; Mismatches 84; Indels 94; Gaps 19;
Qу
          4 LSPLSFIKVSHVRMQGIL---LLVFAKY-----QHL----PYIQILSTKSTPTGLFG 48
            Db
        188 LSP-TYPAAHHFR-SGIIGSGLCVFSKHPIQELTQHIYTLNGYPYMI-------HHG 235
         49 YWGNKGGVN-ICLKLYGYYVSIINCHLPPHISNNYQRLEHFDRILEMQNCEGRDIPNILD 107
Qу
             236 DWFSGKAVGLLVLHLNGMELNAYVTHL--HAEYNRQK----DIYLTHLVGQAWELAQFIH 289
ďŪ
        108 H-----DLIIWFGDMNFRIEDFGLHFVRESIKNRCYGGLWEKDOLSIAKKHDPLL--REF 160
QУ
                290 HTSKKADVVLLCGDLNMHPEDLGCCLLKE-----WTGL------HDAYLETRDF 332
Db
        161 ---OEGRLLFPPTYKFDRNSNDYDTSEKKRKP----AWTDRILWRLKROPCAGPDTPIPP 213
Qу
               Db
         333 KGSEEGNTMVP------KNCY-VSQQELKPFPFGVRIDYVLYK------A 369
        214 ASHFSLSLR-----GYSSHMTYGISDHKPVSGTFDLELKP 248
Qу
             Db
         370 VSGFYISCKSFETTTGFDPHSGTPLSDHEALMATLFVRHSP 410
RESULT 7
US-10-496-011-67
; Sequence 67, Application US/10496011
; GENERAL INFORMATION:
; APPLICANT: BOURGERON, THOMAS
 APPLICANT: JAMAIN, STEPHANE
  APPLICANT: QUACH, HELENE
  APPLICANT: BETANCUR, CATALINA
  APPLICANT: LEBOYER, MARION
  APPLICANT: GILLBERG, CHRISTOPHER
  TITLE OF INVENTION: POLYNUCLEOTIDE AND PROTEIN INVOLVED IN SYNAPTOGENESIS,
  TITLE OF INVENTION: THEREOF, AND THEIR THERAPEUTIC AND DIAGNOSTIC USES
  FILE REFERENCE: 253820US0XPCT
  CURRENT APPLICATION NUMBER: US/10/496,011
  CURRENT FILING DATE: 2004-05-28
  PRIOR APPLICATION NUMBER: PCT/FR02/04134
; PRIOR FILING DATE: 2002-11-28
; PRIOR APPLICATION NUMBER: CA2364106
; PRIOR FILING DATE: 2001-11-30
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; NUMBER OF SEQ ID NOS: 98
; SOFTWARE: PatentIn version 3.3
; SEQ ID NO 67
  LENGTH: 815
   TYPE: PRT
   ORGANISM: Homo sapiens
US-10-496-011-67
 Query Match
                    4.3%; Score 86.5; DB 6; Length 815;
 Best Local Similarity 18.6%; Pred. No. 11;
 Matches 84; Conservative 60; Mismatches 134; Indels 173; Gaps
         5 SPLSFIKVSH------VRMQGILLLVFA-KYQHLPYIQILSTKSTPTGLFGYWGNKG 54
Qу
           257 SCVSLLTLSHYSEGLFOKAIIOSGTALSSWAVNYQPAKYTRILA------DKV 303
        55 GVNI-----PHISNNYQRLEH 87
Qу
           304 GCNMLDTTDMVECLRNKNYKELIQQTITPATYHIAFGPVIDGDVIPDDPQILMEQGEFLN 363
Db
       88 FDRILEMONCEG-RDIPNILDHDLIIWFGDMNFRIEDF-----GLHFVRESIKNRC 137
Qу
           {\tt 364. YDIMLGVNQGEGLKFVDGIVDNEDGVTPNDFDFSVSNFVDNLYGYPEGKDTLRETIK-FM~422}
Db
        138 YGGLWEKDQLSIAKKHDPLLREFQEGRLLFP-----PTYKFDRNSNDYDTSE 184
Qy
           : ::
                   423 YTDWADKENPETRRK--TLVALFTDHQWVAPAVATADLHAQYGSPTYFY----AFYHHCQ 476
Db
       185 KKRKPAWTDRILWRLKROPCAGPDTPIPPASHFSLSLRGYSSHMTYGISDHKPVSGTFDL 244
Qу
                                      : ||:| |
        Db
        245 ELKPLVSAPLIVLMPEDLWTV---ENDMMVS-----YSSTSDFPSSPWDWIGLYKV 292
Qу
           499 ---PMIG------PTELFSCNFSKNDVMLSAVVMTYWTNFAKTGD-PNQPVP----- 540
Db
        293 GLRDV------NDYVSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEF-----L 335
Qу
            541 --QDTKFIHTKPNRFEEVAW---SKYNPKD---QLYLHIGLKPRVRDHYRATKVAFWLEL 592
       336 LCYYSNSLRSVVGISRPFQIPPGSLREDPLG 366
QУ
           :: | :| :| : | |
        593 VPHLHNLNEIFQYVSTTTKVPPPDMTSFPYG 623
RESULT 8
US-10-496-011-3
; Sequence 3, Application US/10496011
; GENERAL INFORMATION:
; APPLICANT: BOURGERON, THOMAS
  APPLICANT: JAMAIN, STEPHANE
; APPLICANT: QUACH, HELENE ; APPLICANT: BETANCUR, CATALINA
; APPLICANT: LEBOYER, MARION
  APPLICANT: GILLBERG, CHRISTOPHER
  TITLE OF INVENTION: POLYNUCLEOTIDE AND PROTEIN INVOLVED IN SYNAPTOGENESIS,
VARIANTS
```

```
TITLE OF INVENTION: THEREOF, AND THEIR THERAPEUTIC AND DIAGNOSTIC USES
; FILE REFERENCE: 253820US0XPCT
  CURRENT APPLICATION NUMBER: US/10/496,011
  CURRENT FILING DATE: 2004-05-28
  PRIOR APPLICATION NUMBER: PCT/FR02/04134
  PRIOR FILING DATE: 2002-11-28
  PRIOR APPLICATION NUMBER: CA2364106
  PRIOR FILING DATE: 2001-11-30
 NUMBER OF SEQ ID NOS: 98
  SOFTWARE: PatentIn version 3.3
; SEO ID NO 3
  LENGTH: 851
   TYPE: PRT
   ORGANISM: Homo sapiens
US-10-496-011-3
                   4.3%; Score 86.5; DB 6; Length 851;
 Query Match
 Best Local Similarity 18.6%; Pred. No. 11;
 Matches 84; Conservative 60; Mismatches 134; Indels 173; Gaps 22;
         5 SPLSFIKVSH-----VRMQGILLLVFA-KYQHLPYIQILSTKSTPTGLFGYWGNKG 54
Qу
          293 SCVSLLTLSHYSEGLFQKAIIQSGTALSSWAVNYQPAKYTRILA------DKV 339
Db
        55 GVNI-----PHISNNYQRLEH 87
QУ
           Db
       340 GCNMLDTTDMVECLRNKNYKELIOOTITPATYHIAFGPVIDGDVIPDDPOILMEOGEFLN 399
        88 FDRILEMONCEG-RDIPNILDHDLIIWFGDMNFRIEDF-----GLHFVRESIKNRC 137
Ov.
           400 YDIMLGVNOGEGLKFVDGIVDNEDGVTPNDFDFSVSNFVDNLYGYPEGKDTLRETIK-FM 458
Db
       138 YGGLWEKDOLSIAKKHDPLLREFOEGRLLFP------PTYKFDRNSNDYDTSE 184
Qу
          459 YTDWADKENPETRRK--TLVALFTDHQWVAPAVATADLHAQYGSPTYFY----AFYHHCQ 512
Db
       185 KKRKPAWTDRILWRLKROPCAGPDTPIPPASHFSLSLRGYSSHMTYGISDHKPVSGTFDL 244
Qу
           : ||:| | : ||
       513 SEMKPSWAD------ 534
Db
       245 ELKPLVSAPLIVLMPEDLWTV---ENDMMVS-----YSSTSDFPSSPWDWIGLYKV 292
Qу
           535 ---PMIG------ PTELFSCNFSKNDVMLSAVVMTYWTNFAKTGD-PNQPVP----- 576
Db
       293 GLRDV-----NDYVSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEF-----L 335
Qy
           577 --ODTKFIHTKPNRFEEVAW---SKYNPKD---OLYLHIGLKPRVRDHYRATKVAFWLEL 628
Db
       336 LCYYSNSLRSVVGISPPFQIPPGSLREDPLG 366
Qу
          :: | ::|| : | |
       629 VPHLHNLNEIFOYVSTTTKVPPPDMTSFPYG 659
Dh
RESULT 9
US-60-643-717-11816
; Sequence 11816, Application US/60643717
; GENERAL INFORMATION:
```

```
; APPLICANT: Abad, Mark S.
  TITLE OF INVENTION: Genes and Uses for Plant Improvement
  FILE REFERENCE: 38-21(53629)A
  CURRENT APPLICATION NUMBER: US/60/643,717
  CURRENT FILING DATE: 2005-01-12
  NUMBER OF SEO ID NOS: 19247
; SEO ID NO 11816
   LENGTH: 321
   TYPE: PRT
   ORGANISM: Candidatus Blochmannia floridanus
US-60-643-717-11816
  Query Match
                         4.2%; Score 85; DB 8; Length 321;
  Best Local Similarity 23.8%; Pred. No. 4.3;
  Matches
          35; Conservative 23; Mismatches
                                              67; Indels 22; Gaps
                                                                         5:
          51 GNKGGVN----ICLKLYGYYVSIINCHLPPHISNNYQRLEHFDRILEMQNCEGRDIPNIL 106
Qу
                     Db
         177 GHKAAVNHVNLLCKKAGAKYVTTLPISVPSHCSIMKKMVSRFQKIIENTVISSPKIPVIN 236
         107 DHDLIIWFGDMNFRIEDFGLHFVRESIKNRCYGGL-WEKDQLSIAKKHDPLLREFQEG-- 163
Qу
                            : |:|:|: : | : | :
             : |: |
                                                        | :: |
         237 NVDVSI-----EQEPQFIRDSLIRQLYNPVRWNEIMQEFINKDIKIVLEMGPGKV 286
Db
         164 -- RLLFPPTYK--- FDRNSNDYDTSEK 185
QУ
               ||: |
                         Db
         287 LTRLIQRSVYSDSLFSLSINDVDSLSK 313
RESULT 10
US-10-450-763-34667
; Sequence 34667, Application US/10450763
; GENERAL INFORMATION:
  APPLICANT: Hyseq, Inc
  TITLE OF INVENTION: NOVEL NUCLEIC ACIDS AND POLYPEPTIDES
  FILE REFERENCE: 790CIP3/US
  CUPRENT APPLICATION NUMBER: US/10/450,763
  CURRENT FILING DATE: 2003-06-11
  PRIOR APPLICATION NUMBER: PCT/US01/08631
  PRIOR FILING DATE: 2001-03-30
  PRIOR APPLICATION NUMBER: 09/540,217
  PRIOR FILING DATE: 2000-03-31
  PRIOR APPLICATION NUMBER: 09/649,167
  PRIOR FILING DATE: 2000-08-23
  NUMBER OF SEQ ID NOS: 60736
  SOFTWARE: Custom
 SEQ ID NO 34667
   LENGTH: 512
   TYPE: PRT
   ORGANISM: Homo sapiens
   FEATURE:
   NAME/KEY: DOMAIN
   LOCATION: (116)..(143)
   OTHER INFORMATION: Mrp family proteins domain identified by eMATRIX,
   OTHER INFORMATION: number BL01215A, p-value=4.964e-11, raw score of 9.75
   FEATURE:
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NAME/KEY: DOMAIN
   LOCATION: (211) . . (302)
   OTHER INFORMATION: ParA family ATPase domain identified by PFam, accession
name
   OTHER INFORMATION: ParA, E-value=0.011, PFam score of -1.9
ï
   FEATURE:
   NAME/KEY: misc feature
   LOCATION: (1)...(512)
   OTHER INFORMATION: Xaa = X or * as defined in Table 2
US-10-450-763-34667
                      4.2%; Score 85; DB 6; Length 512;
 Query Match
 Best Local Similarity 21.6%; Pred. No. 8;
 Matches 68; Conservative 43; Mismatches 124; Indels 80; Gaps
        46 LFGYWGNKGGVNICLKLYGYYVSIINCHLPPHISNNYQRLEHFDRILEMQNCEGRDIPNI 105
Qу
           119 VIGVAAHKGGV-----YKTSVSVHLAQDLAL-----KGLRVLLVEGNDPQGT 160
Db
        106 --LDHDLIIWFGDMNFRIEDFGLHF------VRESIKNRCYGGL-WEKDQLSIAKKHDP 155
Qу
             Db
        161 ASMYHG---WVPDLHIHAEDTLLPFYLGEKDDVTYAIKPTCWPGLDIIPSCLALHRIETE 217
        156 LLREFQEGRLLFPP----TYKFDRNSNDYDTSEKKRKP---AWTDRILWRLKRQPCAGPD 208
Qу
            Db
        218 LMGKFDEGKLPTDPHLMLRLGIETVAHDYDVLVIDSPPNLGIGTINVRWAADVOP---- 272
        209 TPIPPASHFSLSLRGYSSHMTYGISDHKPVSGTFDLE-LKPLVSAPLIVLMPEDLWTVEN 267
QУ
                   273 ------LXIPKQQQNEKYQVPQ------FDQSTIKNIESAKGLDVW--DSWPLQN 313
Db
        268 -DMMVS----YSSTSDFPSSPWDWIGLYKVGLRDVNDYVSYAWVGDSKVSCSDNLNQVYI 322
QУ
            Db
        314 ADGTVAEYNGYHVVFALAGSPKD------ADDTSIYMFYQKVGDNSIDSWKNAGRVFK 365
        323 DI----SNIPTTEDE 333
Qу
              :| | :|:
           Db
        366 DSDKFDANDPILKDQ 380
RESULT 11
US-11-021-951-123
; Sequence 123, Application US/11021951
; GENERAL INFORMATION:
; APPLICANT: HAUPTS, Ulrich
 APPLICANT: KOLTERMANN, Andre
; APPLICANT: SCHEIDIG, Andreas
; APPLICANT: VOTSMEIER, Christian
; APPLICANT: Kettling, Ulrich
; APPLICANT: COCO, Wayne Michael
  TITLE OF INVENTION: New Biological Entities And The Pharmaceutical
  TITLE OF INVENTION: And Diagnostic Use Thereof
; FILE REFERENCE: 04156.0002U5
; CURRENT APPLICATION NUMBER: US/11/021,951
; CURRENT FILING DATE: 2004-12-22
; PRIOR APPLICATION NUMBER: 10/872,198
; PRIOR FILING DATE: 2004-06-18
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PRIOR APPLICATION NUMBER: 60/543,518
  PRIOR FILING DATE: 2004-02-11
  PRIOR APPLICATION NUMBER: 60/524,960
  PRIOR FILING DATE: 2003-11-25
  PRIOR APPLICATION NUMBER: EP 04003058
  PRIOR FILING DATE: 2004-02-11
  PRIOR APPLICATION NUMBER: EP 03025871
  PRIOR FILING DATE: 2003-11-11
  PRIOR APPLICATION NUMBER: EP 03025851
  PRIOR FILING DATE: 2003-11-10
  PRIOR APPLICATION NUMBER: EP 03013819
 PRIOR FILING DATE: 2003-06-18
 NUMBER OF SEQ ID NOS: 191
 SOFTWARE: PatentIn version 3.1
; SEQ ID NO 123
 LENGTH: 735
   TYPE: PRT
  ORGANISM: Homo sapiens
US-11-021-951-123
 Query Match
                     4.2%; Score 85; DB 7; Length 735;
 Best Local Similarity 18.9%; Pred. No. 13;
 Matches 68; Conservative 53; Mismatches 136; Indels 102; Gaps 14;
         62 LYGYYVSIINCHLPPHISNNYQRLEHFDRILEMQNCEGRDIPNILDH-DLIIWFGDMNFR 120
QУ
           19 LLGYYFSDLNFQAPMVVTSS-----TTGDLSIPSSELENIPSENQYFQSAIWSGFIKVK 72
Db
Qу
        121 IEDFGLHFVRESIKNRCYGGLWEKDQLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSND- 179
           Db
         73 KSD--EYTFATSADN--HVTMWVDDQEVINKASNSNKIRLEKGR-LYQIKIQYQRENPTE 127
        180 -----PDTSEKKRKPAWTDRILW------RLKRQPCAGPDTP-----IPPAS 215
Qу
                  Db
        128 KGLDFKLYWTDSQNKKEVISSDNLQLPELKQKSSNSRKKRSTSAGPTVPDRDNDGIPD-- 185
        216 HFSLSLRGY-----SSHMTYGI-----SDHKPVSGTF 242
QУ
           || : |:|
        186 --SLEVEGYTVDVKNKRTFLSPWISNIHEKKGLTKYKSSPEKWSTASDPYSDFEKVTGRI 243
Db
        243 DLELKPLVSAPLIVLMP-----EDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRD 296
Qу
           244 DKNVSPEARHPLVAAYPIVHVDMENIILSKNEDQSTQNTDSQ-----TRT 288
        297 VNDYVSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFQI 355
Qу
           289 ISKNTSTSRTHTSEVHGNAEVHASFFDIGGSVSAG-----FSNSNSSTVAIDHSLSL 340
Db
RESULT 12
US-10-450-763-51703
; Sequence 51.703, Application US/10450763
; GENERAL INFORMATION:
; APPLICANT: Hyseq, Inc
; TITLE OF INVENTION: NOVEL NUCLEIC ACIDS AND POLYPEPTIDES
; FILE REFERENCE: 790CIP3/US
; CURRENT APPLICATION NUMBER: US/10/450,763
```

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CURRENT FILING DATE: 2003-06-11
  PRIOR APPLICATION NUMBER: PCT/US01/08631
  PRIOR FILING DATE: 2001-03-30
  PRIOR APPLICATION NUMBER: 09/540,217
  PRIOR FILING DATE: 2000-03-31
  PRIOR APPLICATION NUMBER: 09/649,167
  PRIOR FILING DATE: 2000-08-23
  NUMBER OF SEO ID NOS: 60736
  SOFTWARE: Custom
; SEQ ID NO 51703
   LENGTH: 1388
   TYPE: PRT
   ORGANISM: Homo sapiens
   FEATURE:
   NAME/KEY: DOMAIN
   LOCATION: (1293)..(1318)
   OTHER INFORMATION: Ubiquitin carboxyl-terminal hydrolases family 2 proteins
   OTHER INFORMATION: domain identified by eMATRIX, accession number BL00972D,
p-value=
   OTHER INFORMATION: 7.750e-19, raw score of 22.55
   FEATURE:
   NAME/KEY: DOMAIN
   LOCATION: (1290)..(1350)
   OTHER INFORMATION: Ubiquitin carboxyl-terminal hydrolase family domain
   OTHER INFORMATION: identified by PFam, accession name UCH-2, E-value=1.1e-
26, PFam
   OTHER INFORMATION: score of 102.0
US-10-450-763-51703
                      4.2%; Score 85; DB 6; Length 1388;
 Query Match
 Best Local Similarity 20.5%; Pred. No. 30;
 Matches 83; Conservative 59; Mismatches 160; Indels 102; Gaps 22;
         32 PYIQILSTKSTPTGLFG--YWGN--KGGVNICLKLY-GYYVSIINCHLPPHISNNYQRLE 86
Qу
            665 PYVELKDSDGRPDWEVAAEAWDNHLRRNRSIIVDLFHGQLRSQVKCKTCGHISVRFDPFN 724
Db
         37 HFDRILEMONCEGRDIPNI-LDHDLIIWFG-DMNFRIEDFGLHFVRESIKNRCYGGLWEK 144
QУ
                725 FLSLPLPMDSYMDLEITVIKLDGTTPVRYGLRLNMDEKYTGL---KKOLRDLC--GL-NS 778
Db
        145 DQLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDYDTSEKKRKPAWTDRILWRLKRQPC 204
QУ
            779 EQILLAEVHDSNIKDFSSS----PST----NGMFTLTTNGDLPKPIFI------PN 820
Db
        205 AGPDTPIPPASHFSLSLRGYSSHMT-----YGISDHKPVSGTFDLELKPLVSAPLI 255
QУ
              821 GMPNTVVPCGTEKNFTNGMVNGHMPSLPDSPFTGYIIAVHRKMMRTELYFLSPQENRPSL 880
Db
        256 VLMPEDLWTVENDMMVS---YSSTSDFPSSPW---DWIG------ 288
QУ
             :: :: :: :: ::
        881 FGMP-----LIVPCTVHTQKKDLYDAVWIQVSWLARPLPPQEASIHAQDRDNCMGY 931
Db
        289 LYKVGLRDV-NDYVSYAWVGD-----SKVSCSDN---LNQVYIDISNIPTTEDEFLLCY 338
QУ
            932 OYPFTLRVVOKDGISCAWCPOYRFCRGCKIDCGEDRAFIGNAYIAVDWHPTA---LHLRY 988
Db
```

```
339 YSNSLRSV-----VGISRPFQIPPGSL-----REDPLGEAQ 369
Qу
            989 QTSQERVVDKHESVEQSRRAQAEPINLDSCLRAFTSEEELGESE 1032
Db
RESULT 13
US-10-450-763-43858
; Sequence 43858, Application US/10450763
; GENERAL INFORMATION:
; APPLICANT: Hyseq, Inc
  TITLE OF INVENTION: NOVEL NUCLEIC ACIDS AND POLYPEPTIDES
; FILE REFERENCE: 790CIP3/US
; CURRENT APPLICATION NUMBER: US/10/450,763
  CURRENT FILING DATE: 2003-06-11
  PRIOR APPLICATION NUMBER: PCT/US01/08631
  PRIOR FILING DATE: 2001-03-30
  PRIOR APPLICATION NUMBER: 09/540,217
  PRIOR FILING DATE: 2000-03-31
  PRIOR APPLICATION NUMBER: 09/649,167
  PRIOR FILING DATE: 2000-08-23
  NUMBER OF SEQ ID NOS: 60736
 SOFTWARE: Custom
; SEQ ID NO 43858
  LENGTH: 716
  TYPE: PRT
  ORGANISM: Homo sapiens
  FEATURE:
  NAME/KEY: DOMAIN
   LOCATION: (235)..(278)
   OTHER INFORMATION: 1 kw MIND INHIBITOR DIVISION CONTROL domain identified by
   OTHER INFORMATION: eMATRIX, accession number DM01756B, p-value=4.822e-15,
raw score of
   OTHER INFORMATION: 22.91
US-10-450-763-43858
 Query Match
                     4.2%; Score 84.5; DB 6; Length 716;
 Best Local Similarity 22.7%; Pred. No. 14;
 Matches 61; Conservative 30; Mismatches 89; Indels 89; Gaps
       138 YGGLW---EKDQLSIAKKHDPLLREFQEGRL-LF----PPTYKF-----DRNSNDYDT 182
Qу
           Db
        427 YGESWNLLRADQRLIFAKSWPRASRYQQGHQDLFILRSDLPSQVFIRDKLMERRNRRTGR 486
        183 SEKKRKPAWTDRIL--WRLKROPCAGPD-----TPIPPASHFSLSLRGYSSHMTY----- 230
Qу
           Db
        487 TEKARIWEVTDRTVRTWIGEAVAAAAADGVTFSVPVTPHTF----RHSYAMHMLYAGIPL 542
        231 ----GISDHKPVSGT------FDLELKPLVSAPLIV-----LMPE----- 260
Qу
               D'n
        543 KVLQSLMGHKSISSTEVYTKVFALD----VAAPARIPKQQQNEKYQVPQFDQSTIKNIES 598
        261 -----DLWTVEN-DMMVS----YSSTSDFPSSPWDWIGLYKVGLRDVNDYVSYAWVGD 308
Qу
                599 AKGLDVWDSWPLQNADGTVAEYNGYHVVFALAGSPKD-----ADDTSIYMFYQKVGD 650
Db
        309 SKVSCSDNLNQVYIDI----SNIPTTEDE 333
Qу
```

```
RESULT 14
US-10-450-763-48706
; Sequence 48706, Application US/10450763
; GENERAL INFORMATION:
; APPLICANT: Hyseq, Inc
  TITLE OF INVENTION: NOVEL NUCLEIC ACIDS AND POLYPEPTIDES
  FILE REFERENCE: 790CIP3/US
  CURRENT APPLICATION NUMBER: US/10/450,763
  CURRENT FILING DATE: 2003-06-11
; PRIOR APPLICATION NUMBER: PCT/US01/08631
  PRIOR FILING DATE: 2001-03-30
  PRIOR APPLICATION NUMBER: 09/540,217
  PRIOR FILING DATE: 2000-03-31
  PRIOR APPLICATION NUMBER: 09/649,167
  PRIOR FILING DATE: 2000-08-23
; NUMBER OF SEQ ID NOS: 60736
  SOFTWARE: Custom
; SEQ ID NO 48706
   LENGTH: 731
   TYPE: PRT
   ORGANISM: Homo sapiens
   FEATURE:
   NAME/KEY: DOMAIN
   LOCATION: (620)..(635)
   OTHER INFORMATION: GLYCOSYL HYDROLASE FAMILY 1 SIGNATURE domain identified
by
   OTHER INFORMATION: eMATRIX, accession number PR00131A, p-value=1.563e-11,
raw score of
   OTHER INFORMATION: 12.98
   FEATURE:
   NAME/KEY: DOMAIN
   LOCATION: (531)..(719)
   OTHER INFORMATION: Glycosyl hydrolase family 1 domain identified by PFam,
   OTHER INFORMATION: accession name Glyco hydro 1, E-value=1.3e-43, PFam score
of 155.2
US-10-450-763-48706
                        4.2%; Score 84.5; DB 6; Length 731;
 Query Match
 Best Local Similarity 18.5%; Pred. No. 14;
 Matches 74; Conservative 48; Mismatches 119; Indels 159; Gaps 19;
QУ
          54 GGVNICLK-LYGYYVSIINC-HLP-------PHISNNYQRLEHF 88
             11 :: : 1
         343 GNINNALKYAKGEFVSIFDCDHVPTRSFLQMTMGWFLKEKQLAMMQTPH---HFFSPDPF 399
Db
          89 DRILEMONCEGRDIPNILDHDLIIWFG-----DMNFRIEDFGLHFVRESIKNRCYGGLWE 143
Qу
             400 ERNLG----RFRKTPN----EGTLFYGLRQSWEFNAR----GLELVKEGRAQACVSAGNT 447
Db
         144 KDQLSIAKKHDPLLR--EFQEGRLLFPPTYKFDRNSNDYDTSEKKRKPAWTDRILWRLKR 201
Qу
               : :|| || :|
Db
         448 GALMGLAKL---LLKPLEGONHRMLL------RWEAKLAALVRL 482
         202 OPCAGPDTPIPPAS-----HFSLSLRGYSSHMTYGISDHKPVSGTFDLELKPLVSA 252
Qу
```

```
483 QENTAQPVPFAPNNARPLTLEDDRLSCTVRGYNFAITFSKMSGKPTSW------QLVAS 535
Db
         253 PLIVLM------PEDLWTV-----ENDMMVSYSSTSDFPS- 281
Qу
                                 : '
         536 ALATKIAHEVNPQNQVGCMLAGGNFYPYSCKPEDVWAALEKDRENLFFIDVQARGTYPAY 595
Db
         282 -----SPWDWIGLYKVGLRDVNDYVSYAWVGDSKVSCSDNLNQVYIDISNI 327
Oy
                         :| | | : | : | : | : | |
         596 SARVFREKGVTINKAPGD----- 642
Db
         328 PTTEDEFLLCYYSNSLRSVVGISRPF-OIPPGSLREDPLG 366
QУ
                       | :: | : |: |: | | | | |
         643 -----SSAANVVKSLRNPYLQVSDWGWGIDPLG 670
Db
RESULT 15
PCT-US04-17965-2224
; Sequence 2224, Application PC/TUS0417965
; GENERAL INFORMATION:
; APPLICANT: ARBORGEN, LLC
 APPLICANT: BLOKSBERG, LEONARD N.
 APPLICANT: BRYANT, CATHERINE
; APPLICANT: CONNETT, MARIE B.
; APPLICANT: EMERSON, SARAH JANE
; APPLICANT: FROST, MICHAEL J.
; APPLICANT: FORSTER, RICHARD LLEWELLYN SYDNEY
; APPLICANT: GRIGOR, MURRAY
; APPLICANT: HIGGINS, COLLEEN
; APPLICANT: LASHAM, ANNETTE
  APPLICANT: LUND, STEVEN TROY APPLICANT: MAGUSIN, ANDREAS
  APPLICANT: PHILLIPS, JONATHAN
  APPLICANT: PUTHIGAE, SATHIAH
  APPLICANT: VEERAKONE, STELLA
 APPLICANT: WESTWOOD, CLAIRE
 APPLICANT: GAUSE, KATRINA
 APPLICANT: WOOD, MARION
APPLICANT: ROTTMAN, WILLIAM
; APPLICANT: HAVUKKALA, ILKKA
; TITLE OF INVENTION: TRANSCRIPTION FACTORS
; FILE REFERENCE: 044463-0296
; CURRENT APPLICATION NUMBER: PCT/US04/17965
; CURRENT FILING DATE: 2004-06-07
  PRIOR APPLICATION NUMBER: 60/476,189
; PRIOR FILING DATE: 2003-06-06
; NUMBER OF SEQ ID NOS: 3679
 SOFTWARE: PatentIn Ver. 3.2
; SEQ ID NO 2224
   LENGTH: 380
   TYPE: PRT
   ORGANISM: Pinus radiata
PCT-UC04-17965-2224
 Query Match
                       4.2%; Score 84; DB 1; Length 380;
 Best Local Similarity 21.4%; Pred. No. 6.7;
 Matches 65; Conservative 37; Mismatches 96; Indels 106; Gaps
```

Qу	92	LEMQNCEGRDIPNILDHDLIIWFGDMNFRIEDFGLHFVRESIKNR-CYGGL	141
Db	100	LEVQKRRIYDITNVLEGIGLIEKRLKNRICWKGLSVSRPGEVE	142
Qy	142	wekdqlsiak-khdpllrefqeg-rllfpptykfdrnsndydtsekkrkpaw-	191
Db	143	DEATVLQAEVDSLNLEECKLDDCIRDMQERLRILSEDDRNRRWL	186
Qy	192	TDRILWRLKRQPCAGPDT	229
Db	187	YVTDEDIKKLPCFQNDTLIAIKAPHGTTLEVPDPDEAVEYPQRRYQILLRST	238
Qу	230	YGISDHKPVSGTFDLELKPLVSAPLIV-LMPEDLWTVENDMMVSYSSTSDFP	280
Db	239	MGPIDVYLVS-QFEENIEEMNPVDLATELVPSGICPAEGVTISSVQEGATFVEMECQGHE	297
Qy	281	SSPWDWI-GLYKVGLRDVNDYVSYAWVGDSKVSCSDNLNQVYIDISNI:	327
Db	298	TRQPCTELTSPQDSAGGIMRIVPSDANIDADYWLLSDSGVGITDMWRTETNSTWDEVIEL	357
Qy	328	PTTE 331	
Db	3,58	NPTD 361	,

Search completed: February 10, 2005, 18:02:13
Job time: 25 secs

GenCore version 5.1.6 Copyright (c) 1993 - 2005 Compugen Ltd.

OM protein - protein search, using sw model

Run on: February 10, 2005, 17:03:41; Search time 166 Seconds

(without alignments)

866.716 Million cell updates/sec

Title: US-09-892-287-1

Perfect score: 2020

Sequence: 1 MDVLSPLSFIKVSHVRMQGI......FQIPPGSLREDPLGEAQPQI 372

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 2105692 seqs, 386760381 residues

Total number of hits satisfying chosen parameters: 2105692

Minimum D3 seq length: 0

Maximum DB seg length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : A Geneseq 16Dec04:*

1: geneseqp1980s:*

2: geneseqp1990s:*

3: geneseqp2000s:*

4: geneseqp2001s:*

5: geneseqp2002s:*

6: geneseqp2003as:*

7: geneseqp2003bs:*

8: geneseqp2004s:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result		% Query				
No.	Score		Length	DB	ID	Description
1	2020	100.0	372	2	AAW97094	Aaw97094 Phosphati
2	2020	100.0	448	6	AAE36070	Aae36070 Human pho
3	2012	99.6	478	6	AAE36067	Aae36067 Human pho
4	2010	99.5	372	6	ABO07198	Abo07198 Human p53
5	2010	99.5	372	6	AAE36069	Aae36069 Human pho
6	2010	99.5	448	6	AAE36068	Aae36068 Human pho
7	2010	99.5	448	6	ABO53040	Abo53040 Human put
8	1025	50.7	240	8	ADN61442	Adn61442 Human KPP
9	872.5	43.2	1001	5	AAU98903	Aau98903 Rat inosi

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638 8 ADR08504
10
     870.5
            43.1
                                                       Adr08504 Human pro
11
     868.5
             43.0
                     639 7
                             ADB65114
                                                       Adb65114 Human pro
12
     868.5
             43.0
                     639
                         7
                             ADM04740
                                                       Adm04740 Human pro
                                                       Aau98904 Human ino
13
     868.5
            43.0
                    1006 5
                            AAU98904
14
     868.5
            43.0
                    1006
                          6
                             ABU54583
                                                       Abu54583 Human NOV
15
     831.5
            41.2
                     751
                         5
                            AAU98902
                                                       Aau98902 Human ino
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            41.2
                    1056
                          6
                            AAE36066
                                                       Aae36066 Human pho
17
    683.5
            33.8
                     279
                            AAG73981
                                                       Aaq73981 Human col
18
    637.5
            31.6
                     254
                            ABR01000
                                                       Abr01000 Human gen
                          6
            31.6
                     254 6
                                                       Ada98192 Human sec
19
    637.5
                            ADA98192
20
    637.5
            31.6
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                                                       Aab27797 Human sec
                     162
                                                       Abp43025 Human ova
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    541.5
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                         5
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    507.5
            25.1
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22
                            ABB64662
                     397 7
23
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                                                       Adj68405 Human hea
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    494.5
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                     381 3
                            AAB27845
                                                       Aab27845 Sequence
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    460.5
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                     357
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                                                       Abb62704 Drosophil
                          4
26
    460.5
            22.8
                     357
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27
    432.5
            21.4
                     749
                         4
                             AAB95181
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28
                                                       Aab27846 Protein f
       428
            21.2
                     382
                          3
                             AAB27846
29
            21.0
                     776
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                         8
                                                       Adt49828 Murine IN
30
    367.5
            18.2
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                            ADJ69678
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       328
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       328
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                            ABU08111
                                                       Abu08111 Human kin
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       328
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            16.2
                                                       Adr14671 Human NF-
39
                   1496 8
                            ADR14671
    327.5
            16.2
                            ABM80825
                                                       Abm80825 Tumour-as
40
                    1496
                         8
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            16.2
                    1510
                          5
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                                                       Abb76319 Human pro
42
      326
            16.1
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                         7
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                                                       Add46315 Rat Prote
43
       326
                                                       Ade63137 Rat Prote
            16.1
                    1248
                         7
                             ADE63137
44
     316.5
            15.7
                    1114
                             AAU76370
                                                       Aau76370 Drosophil
45
    316.5
            15.7
                    1218
                             ABB62412
                                                       Abb62412 Drosophil
```

ALIGNMENTS

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     AAW97094 standard; protein; 372 AA.
ID
XX
AC
     AAW97094;
XX
DT
     28-APR-1999 (first entry)
XX
DE
     Phosphatidylinositol 4, 5-bisphosphate 5-phosphatase.
XX
KW
     Human; phosphatidylinositol 4,5-bisphosphate 5-phosphatase; PBPP;
     Incyte clone 638789; antagonist; immune disorder; cancer;
KW
KW
     neuronal disorder; human tubby homologue.
XX
OS
     Homo sapiens.
XX
```

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Location/Qualifiers
FΗ
     Key
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FT
                     /note= "potential phosphorylation site"
FT
     Modified-site
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                     /note= "potential phosphorylation site"
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FΤ
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FT
FT
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XX
    WO9900507-A1.
PN
XX
PD
     07-JAN-1999.
XX
PF
     26-JUN-1998;
                    98WO-US013399.
XX
PR
     27-JUN-1997;
                    97US-00884681.
XX
PΑ
     (INCY-) INCYTE PHARM INC.
XX
ΡI
     Hillman JL, Lal P, Corley NC, Shah P;
XX
DR
     WPI; 1999-095752/08.
DR
     N-PSDB; AAX15254.
XX
PT
     Phosphatidylinositol 4,5-bisphosphate 5-phosphatase - used for treating
PT
     immune disorders, cancers, and neuronal disorders.
XX
PS
     Claim 1; Fig 1A-G; 81pp; English.
XX
CC
     The present sequence encodes a human phosphatidylinositol 4,5-
     bisphosphate 5-phosphatase (PBPP) protein. The PBPP polynucleotide was
CC
CC
     first identified in Incyte clone 638789 from the breast cancer cDNA
CC
     library BRSTNOTO3. Antagonists of the PBPP protein can be used in the
     treatment or prevention of an immune disorder, a cancer, or a neuronal
CC
CC
     disorder. The PBPP polynucleotide can be used for the detection of
CC
     polynucleotides encoding human tubby homologue. The immune disorders that
CC
     can be treated include AIDS, Addison's disease, adult respiratory
CC
     distress syndrome, allergies, anaemia, asthma, atherosclerosis, Crohn's
CC
     disease, ulcerative colitis, atopic dermatitis, gout, Grave's disease,
CC
     irritable bowel syndrome, lupus erythematosus, multiple sclerosis,
```

```
CC
    scleroderma, and autoimmune thyroditis. Also, complications of cancer,
    haemodialysis, extracorporeal circulation, infection, and trauma can be
CC
CC
    treated using the antagonist. The neuronal disorders that can be treated
    include Alzheimer's disease, amnesia, catatonia, amyotrophic lateral
CC
    sclerosis, dementia, depression, Down's syndrome, epilepsy, Huntington's
CC
CC
    disease, multiple sclerosis, neurofibromatosis, Parkinson's disease,
    paranoid psychoses, schizophrenia and Tourette's syndrome
CC
XX
SO
    Sequence 372 AA;
 Query Match
                      100.0%;
                             Score 2020; DB 2; Length 372;
                      100.0%;
 Best Local Similarity
                             Pred. No. 5.9e-204;
                           0; Mismatches
 Matches 372; Conservative
                                              Indels
                                           0;
                                                       0;
                                                          Gaps
                                                                 0;
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Qу
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Qу
            Db
        121 IEDFGLHFVRESIKNRCYGGLWEKDOLSIAKKHDPLLREFOEGRLLFPPTYKFDRNSNDY 180
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Qу
            Db
        301 VSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFQIPPGSL 360
        361 REDPLGEAOPOI 372
QУ
            361 REDPLGEAQPQI 372
Db
RESULT 2
AAE36070
ID
    AAE36070 standard; protein; 448 AA.
XX
AC
    AAE36070;
XX
DT
    26-JUN-2003 (first entry)
XX
DE
    Human phosphatidylinositol biphosphate (PIB) protein #5.
XX
KW
    Human; p53 pathway; phosphatidylinositol biphosphate; PIB; therapeutic;
    cancer; apoptosis; cell proliferation; cytostatic.
KW
XX
```

myasthenia gravis, osteoarthritis, osteoporosis, rheumatoid arthritis,

CC

```
os
     Homo sapiens.
XX
                    Location/Qualifiers
FΗ
     Key
                    12. .326
FT
     Domain
FT
                    /note= "Inositol polyphosphate phospatase family,
                    catalytic (IPPC) domain"
FT
XX
ΡN
     WO200299125-A1.
XX
     12-DEC-2002.
ΡD
XX
     03-JUN-2002; 2002WO-US017526.
PF
XX
PR
     05-JUN-2001; 2001US-0296076P.
     10-CCT-2001; 2001US-0328605P.
PR
     15-FEB-2002; 2002US-0357253P.
PR
XX
PA
     (EXEL-) EXELIXIS INC.
XX
PΙ
     Friedman L, Plowman GD, Belvin M, Francis-Lang H, Li D, Funke RP;
XX
DR
     WPI; 2003-167348/16.
XX
PT
     Identifying candidate p53 pathway-modulating agents, useful as
PT
     therapeutic targets for disorders associated with defective p53 function,
PT
     comprises screening for agents that modulate phosphatidylinositol
PT
    biphosphate activity.
XX
PS
     Claim 13; Page 133-135; 139pp; English.
XX
CC
     The present invention relates to a method of identifying candidate p53
CC
     pathway-modulating agents. The method involves screening for agents that
CC
    modulate the activity of phosphatidylinositol biphosphate (PIB). The
CC
    methods are useful for identifying candidate p53 pathway-modulating
CC
     agents used as therapeutic targets for disorders associated with
CC
     defective p53 function. They are also useful for modulating p53 pathway
CC
     in a mammalian cell or for diagnosing or treating a disease associated
CC
     with defective p53 function, e.g. cancers such as breast cancer, colon
     cancer, lung cancer or ovarian cancer. Sequences of the invention are
CC
     useful for identifying and testing agents that modulate PIB function. The
CC
     animal models are useful for in vivo assays to test the activity of
CC
CC
     candidate p53-modulating agents or to assess the role of PIB in a p53
CC
     pathway process such as apoptosis or cell proliferation. The present
CC
     sequence is human PIB protein
XX
SO
     Sequence 448 AA;
  Query Match
                         100.0%; Score 2020; DB 6; Length 448;
  Best Local Similarity
                         100.0%; Pred. No. 7.8e-204;
                                                              0; Gaps
  Matches 372; Conservative
                               0; Mismatches
                                                0; Indels
                                                                          0;
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QУ
            317 TFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVNDY 376
Db
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Qу
            377 VSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFQIPPGSL 436
Db
        361 REDPLGEAQPQI 372
Qy
           437 REDPLGEAQPQI 448
Db
RESULT 3
AAE36067
TD
    AAE36067 standard; protein; 478 AA.
XX
AC
    AAE36067;
XX
DT
    26-JUN-2003 (first entry)
XX
DE
    Human phosphatidylinositol biphosphate (PIB) protein #2.
XX
ΚW
    Human; p53 pathway; phosphatidylinositol biphosphate; PIB; therapeutic;
KW
    cancer; apoptosis; cell proliferation; cytostatic.
XX
os
    Homo sapiens.
XX
FH
    Key
                  Location/Qualifiers
FT
    Misc-difference 56
FT
                  /label= Unknown
XX
ΡИ
    WO200299125-A1.
XX
PD
    12-DEC-2002.
XX
PF
    03-JUN-2002: 2002WO-US017526.
XX
PR
    05-JUN-2001; 2001US-0296076P.
Pĸ
    10-OCT-2001; 2001US-0328605P.
    15-FEE-2002; 2002US-0357253P.
PR
XX
PA
    (EXEL-) EXELIXIS INC.
XX
ΡI
    Friedman L, Plowman GD,
                          Belvin M, Francis-Lang H, Li D, Funke RP;
XX
    WPI; 2003-167348/16.
DR
```

```
XX
PT
     Identifying candidate p53 pathway-modulating agents, useful as
     therapeutic targets for disorders associated with defective p53 function,
PT
     comprises screening for agents that modulate phosphatidylinositol
PT
PT
     biphosphate activity.
XX
PS
     Claim 13; Page 128-129; 139pp; English.
XX
     The present invention relates to a method of identifying candidate p53
CC
     pathway-modulating agents. The method involves screening for agents that
CC
     modulate the activity of phosphatidylinositol biphosphate (PIB). The
CC
CC
     methods are useful for identifying candidate p53 pathway-modulating
CC
     agents used as therapeutic targets for disorders associated with
     defective p53 function. They are also useful for modulating p53 pathway
CC
CC
     in a mammalian cell or for diagnosing or treating a disease associated
     with defective p53 function, e.g. cancers such as breast cancer, colon
CC
CC
     cancer, lung cancer or ovarian cancer. Sequences of the invention are
     useful for identifying and testing agents that modulate PIB function. The
CC
CC
     animal models are useful for in vivo assays to test the activity of
CC
     candidate p53-modulating agents or to assess the role of PIB in a p53
CC
     pathway process such as apoptosis or cell proliferation. The present
CC
     sequence is human PIB protein
ХX
SO
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                          99.6%;
                                  Score 2012; DB 6; Length 478;
  Best Local Similarity
                          99.7%;
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  Matches 371; Conservative
                                 0;
                                    Mismatches
                                                   1; Indels
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          Db
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       361 REDPLGEAQPQI 372
Qy
          467 REDPLGEAQPQI 478
Db
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Gaps

0;

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     ABO07198 standard; protein; 372 AA.
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XX
AC
     ABO07198;
XX
     13-AUG-2003 (first entry)
DT
XX
     Human p53 modifying protein, SEQ ID 158.
DΕ
XX
KW
     Human; p53 modifier; cytostatic; cancer; cytostatic; antiangiogenic;
     antiapoptotic; p53 pathway; breast cancer; colon cancer; kidney cancer;
KW
     lung cancer; ovarian cancer; angiogenesis; cell cycle;
KW
KW
     apoptotic disorder; cell proliferation disorder.
XX
os
     Homo sapiens.
XX
PN
     WO200299122-A1.
XX
     12-DEC-2002.
PD
XX
     03-JUN-2002; 2002WO-US017382.
PF
XX
PR
     05-JUN-2001; 2001US-0296076P.
PR
     10-OCT-2001; 2001US-0328605P.
PR
     15-FEB-2002; 2002US-0357253P.
XX
PA
     (EXEL-) EXELIXIS INC.
XX
PΙ
     Friedman L, Plowman GD, Belvin M, Francis-Lang H, Li D, Funke RP;
XX
DR
     WPI; 2003-156859/15.
DR
     N-PSDB; ACD13373.
XX
PT
     Identifying modulators of the p53 pathway for use in treating apoptotic
PT
     or cell proliferation disorders, comprises screening for agents that
PT
     modulate activity of a human ortholog of genes that modify the p53
PT
     pathway in Drosophila.
XX
PS
     Example 2; Page 474-475; 678pp; English.
XX
     The invention relates to identifying (M1) a candidate p53 pathway
CC
     modulating agent, by contacting an assay system comprising a purified HM
CC
CC
     polypeptide (human orthologue of genes that modify the p53 pathway in
     Drosophila) or nucleic acid with a test agent under conditions, where but
CC
CC
     for the presence of the test agent, the system provides a reference
CC
     activity, and detecting a test agent-biased activity of the assay system.
     Also included are modulating (M2) a p53 pathway of a cell (comprising
CC
CC
     contacting a cell defective in p53 function with a candidate mcdulator
CC
     that specifically binds to a HM polypeptide comprising an HM amino acid
     sequence, where p53 function is restored), modulating (M3) a p53 pathway
CC
     in a mammalian cell (comprising contacting the cell with an agent that
CC
     specifically binds an HM polypeptide or nucleic acid) and diagnosing (M4)
CC
     a disease in a patient (comprising: (a) obtaining a biological sample
CC
     from the patient; (b) contacting the sample with a probe for HM
CC
CC
     expression; (c) comparing the results with a control; and (d) determining
```

```
CC
    whether the comparison indicates a likelihood disease). (M1) is useful
    for identifying modulators of the p53 pathway. A probe for HM expression
CC
    is useful for diagnosing breast, colon, kidney, lung and ovarian cancer,
CC
    in a patient, where the cancer has greater than 25 % expression level.
CC
    Modulators identified by (M1) are useful in a variety of diagnostic and
CC
    therapeutic applications, where disease or disorder prognosis is related
CC
    to defects in the p53 pathway, such as, angiogenesis, apoptotic or cell
CC
    proliferation disorders (e.g. cancer). Another two new methods (M2 and
CC
CC
    M3) are useful for modulating the p53 pathway of a cell, thus restoring
    the p53 function of the cell, so that the cell undergoes normal
CC
    proliferation or progression through the cell cycle. (M2) and (M3) are
CC
CC
    also useful for treating defects in the p53 pathway such as angiogenic,
CC
    apoptotic or cell proliferation disorders. The present sequence
CC
    represents a human p53 pathway modifying protein
XX
SO
    Sequence 372 AA;
 Query Match
                      99.5%;
                             Score 2010; DB 6; Length 372;
 Best Local Similarity
                      99.5%;
                             Pred. No. 6.7e-203;
 Matches 370; Conservative
                            0; Mismatches
                                            2;
                                               Indels
                                                                  0;
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Qу
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Qу
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Qу
            181 DTSEKKRKPAWTDRILWRLKROPCAGPDTPIPPASHFSLSLRGYSSHMTYGISDHKPVSG 240
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Qу
            241 TFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVNDY 300
Db
        301 VSYAWVGDSKVSCSDNLNOVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFOIPPGSL 360
Oy
            301 VSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYRNSLRSVVGIRRPFQIPPGSL 360
Db
        361 REDPLGEAOPOI 372
Qy
            361 REDPLGEAQPQI 372
Db
RESULT 5
    AAE36069 standard; protein; 372 AA.
XX
AC
    AAE36069;
XX
    25-JUN-2003 (first entry)
DT
```

```
XX
DE
     Human phosphatidylinositol biphosphate (PIB) protein #4.
XX
     Human; p53 pathway; phosphatidylinositol biphosphate; PIB; therapeutic;
KW
     cancer; apoptosis; cell proliferation; cytostatic.
KW
XX
os
     Homo sapiens.
XX
     WO200299125-A1.
PN
XX
PD
     12-DEC-2002.
XX
PF
     03-JUN-2002; 2002WO-US017526.
XX
PR
     05-JUN-2001; 2001US-0296076P.
     10-OCT-2001; 2001US-0328605P.
PR
     15-FEB-2002; 2002US-0357253P.
PR
XX
PΑ
     (EXEL-) EXELIXIS INC.
XX
PΙ
     Friedman L,
                 Plowman GD, Belvin M, Francis-Lang H, Li D, Funke RP;
XX
DR
     WPI; 2003-167348/16.
XX
РΤ
     Identifying candidate p53 pathway-modulating agents, useful as
PT
     therapeutic targets for disorders associated with defective p53 function,
TС
     comprises screening for agents that modulate phosphatidylinositol
PT
     biphosphate activity.
XX
PS
     Claim 13; Page 131-133; 139pp; English.
XX
CC
     The present invention relates to a method of identifying candidate p53
CC
     pathway-modulating agents. The method involves screening for agents that
CC
     modulate the activity of phosphatidylinositol biphosphate (PIB). The
     methods are useful for identifying candidate p53 pathway-modulating
CC
CC
     agents used as therapeutic targets for disorders associated with
CC
     defective p53 function. They are also useful for modulating p53 pathway
CC
     in a mammalian cell or for diagnosing or treating a disease associated
     with defective p53 function, e.g. cancers such as breast cancer, colon
CC
CC
     cancer, lung cancer or ovarian cancer. Sequences of the invention are
CC
     useful for identifying and testing agents that modulate PIB function. The
CC
     animal models are useful for in vivo assays to test the activity of
CC
     candidate p53-modulating agents or to assess the role of PIB in a p53
CC'
     pathway process such as apoptosis or cell proliferation. The present
CC
     sequence is human PIB protein
XX
SQ
     Sequence 372 AA;
  Query Match
                         99.5%;
                                 Score 2010; DB 6;
                                                     Length 372;
                         99.5%;
                                 Pred. No. 6.7e-203;
  Best Local Similarity
 Matches 370; Conservative
                                0; Mismatches
                                                  2;
                                                      Indels
                                                                            0;
            1 MDVLSPLSFIKVSHVRMQGILLLVFAKYQHLPYIQILSTKSTPTGLFGYWGNKGGVNICL 60
Qу
              DЪ
            1 MDVLSPLSFIKVSHVRMOGILLLVFAKYOHLPYIOILSTKSTPTGLFGYWGNKGGVNICL 60
Qу
           61 KLYGYYVSIINCHLPPHISNNYQRLEHFDRILEMONCEGRDIPNILDHDLIIWFGDMNFR 120
```

```
Db
         61 KLYGYYVSIINCHLPPHISNNYORLEHFDRILEMONCEGRDIPNILDHDLIIWFGDMNFR 120
        121 IEDFGLHFVRESIKNRCYGGLWEKDQLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDY 180
Qy
            Db
        121 IEDFGLHFVRESIKNRCYGGLWEKDQLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDY 180
        181 DTSEKKRKPAWTDRILWRLKROPCAGPDTPIPPASHFSLSLRGYSSHMTYGISDHKPVSG 240
Qу
            181 DTSEKKRKPAWTDRILWRLKROPCAGPDTPIPPASHFSLSLRGYSSHMTYGISDHKPVSG 240
Db
        241 TFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVNDY 300
Qу
            Db
        241 TFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVNDY 300
Qy
        301 VSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFQIPPGSL 360
            Db
        301 VSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYRNSLRSVVGIRRPFQIPPGSL 360
        361 REDPLGEAQPQI 372
Qу
            Db
        361 REDPLGEAQPQI 372
RESULT 6
    AAE36068 standard; protein; 448 AA.
XX
AC
    AAEC6068;
XX
DT
    26-J(JN-2003 (first entry)
XX
DE
    Human phosphatidylinositol biphosphate (PIB) protein #3.
XX
KW
    Human; p53 pathway; phosphatidylinositol biphosphate; PIB; therapeutic;
KW
    cancer; apoptosis; cell proliferation; cytostatic.
XX
os
    Homo sapiens.
XX
PN
    WO200299125-A1.
XX
PD
    12-DEC-2002.
XX
PF
    03-JUN-2002; 2002WO-US017526.
XX
    05-JUN-2001; 2001US-0296076P.
PR
    10-OCT-2001; 2001US-0328605P.
PR
    15-FEB-2002; 2002US-0357253P.
PR
XX
PF.
    (EXEL-) EXELIXIS INC.
XX
    Friedman L, Plowman GD, Belvin M, Francis-Lang H, Li D, Funke RP;
_{
m PI}
XX
DR
    WPI; 2003-167348/16.
XX
PT
    Identifying candidate p53 pathway-modulating agents, useful as
PT
    therapeutic targets for disorders associated with defective p53 function,
```

comprises screening for agents that modulate phosphatidylinositol PT PTbiphosphate activity. ХX Claim 13; Page 130-131; 139pp; English. PS XX The present invention relates to a method of identifying candidate p53 CC pathway-modulating agents. The method involves screening for agents that CC modulate the activity of phosphatidylinositol biphosphate (PIB). The CC methods are useful for identifying candidate p53 pathway-modulating CC agents used as therapeutic targets for disorders associated with CC defective p53 function. They are also useful for modulating p53 pathway CC CC in a mammalian cell or for diagnosing or treating a disease associated with defective p53 function, e.g. cancers such as breast cancer, colon CC cancer, lung cancer or ovarian cancer. Sequences of the invention are CC useful for identifying and testing agents that modulate PIB function. The CCCC animal models are useful for in vivo assays to test the activity of CC candidate p53-modulating agents or to assess the role of PIB in a p53 pathway process such as apoptosis or cell proliferation. The present · CC CC sequence is human PIB protein XX SQ Sequence 448 AA; Query Match 99.5%; Score 2010; DB 6; Length 448; 99.5%; Pred. No. 8.9e-203; Best Local Similarity 0; Mismatches Matches 370; Conservative 2; Indels 0; Gaps 0; 1 MDVLSPLSFIKVSHVRMOGILLLVFAKYOHLPYIQILSTKSTPTGLFGYWGNKGGVNICL 60 QУ 77 MDVLSPLSFIKVSHVRMOGILLLVFAKYOHLPYIOILSTKSTPTGLFGYWGNKGGVNICL 136 Db 61 KLYGYYVSIINCHLPPHISNNYQRLEHFDRILEMQNCEGRDIPNILDHDLIIWFGDMNFR 120 Qу 137 KLYGYYVSIINCHLPPHISNNYORLEHFDRILEMONCEGRDIPNILDHDLIIWFGDMNFR 196 Db 121 IEDFGLHFVRESIKNRCYGGLWEKDQLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDY 180 QУ 197 IEDFGLHFVRESIKNRCYGGLWEKDQLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDY 256 Db 181 DTSEKKRKPAWTDRILWRLKRQPCAGPDTPIPPASHFSLSLRGYSSHMTYGISDHKPVSG 240 Qу 257 DTSEKKRKPAWTDRILWRLKROPCAGPDTPIPPASHFSLSLRGYSSHMTYGISDHKPVSG 316 Db 241 TFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVNDY 300 Qν 317 TFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVNDY 376 Db 30 VSYAWVGDSKVSCSDNLNOVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFOIPPGSL 360: Qу 377 VSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYRNSLRSVVGIRRPFQIPPGSL 436 Db 361 REDPLGEAQPQI 372 Qу

Db

437 REDPLGEAOPOI 448

```
ABO53040 standard; protein; 448 AA.
ID
XX
AC
     ABO53040;
XX
DT
     10-OCT-2003 (first entry)
XX
     Human putative spliceosome associated protein (SAP) #16.
DE
XX
     Human; SAP; spliceosome associated protein; ribonucleoprotein;
KW
     RNP complex; RNA affinity substrate; RNP assembly sequence;
KW
     spliceosomal complex; hnRNP complex; mRNA export complex;
KW
     mRNA localisation complex; RNA editing complex; intron complex;
KW
     H complex; telomerase complex; fragile X protein complex;
KW
KW.
     reverse transcriptase complex; gene splicing complex.
XX
os
     Homo sapiens.
XX
PN
     US2003068803-A1.
XX
PD
     10-APR-2003.
XX
PF
     14-JAN-2002; 2002US-00047991.
XX
PR
     12-JAN-2001; 2001US-0261521P.
XX
PΑ
     (REED/) REED R.
PΑ
     (ZHOU/) ZHOU Z.
XX
PΙ
     Reed R,
              Zhou Z;
XX
     WPI; 2003-540885/51.
DR
XX
PT
     Isolating ribonucleoprotein complex, by contacting RNA affinity substrate
     having ribonucleoprotein assembly sequence and affinity tag, with protein
PT
     mixture, subjecting complex formed to chromatography, affinity selection.
PT
XX
     Claim 24; Page; 39pp; English.
PS
XX
CC
     The invention relates to forming (M1) an isolated ribonucleoprotein (RNP)
     complex (C), involves contacting an RNA affinity substrate (S) comprising
CC
     an RNP assembly sequence (AS) and an affinity tag, with a protein mixture
CC
     to permit formation of (C) on AS, subjecting (C) to chromatographic
CC
CC
     separation, and subjecting (C) to affinity selection, where the affinity
CC
     tag (e.g. bacteriophage MS2 coat protein in a fusion protein with E. coli
CC
     maltose binding protein) binds to an affinity matrix. Also included are
CC
     an isolated spliceosome preparation (isolated by (M1)), a RNA comprising
CC
     an RNP complex binding site and at least one phage coat protein
CC
     recognition site, a nucleic acid encoding the RNA, and treating (M2) a
CC
     subject having a disorder associated with abnormal RNP complexes (by
CC
     obtaining a sample of cells from a subject, purifying RNP complexes from
CC
     the cells of the subject by (M1), determining the presence in the
     purified RNP complexes of one or more proteins, and normalising the
CC
     amount of RNPs in the subject. (M1) is useful for forming an isolated RNP
CC
     complex selected from a spliceosomal complex (selected from E, A, B and C
CC
     complex), an hnRNP complex, an mRNA export complex, an mRNA localisation
CC
     complex, an RNA editing complex, an intron complex, or an H complex. (M1)
CC
     is useful in a diagnostic assay for determining whether a subject has
CC
```

```
CC
     disorder associated with abnormal RNP complexes. (M1) is useful for
 CC
     forming an isolated RNP complex such as a telomerase complex, a fragile X
 CC
     protein complex, a reverse transcriptase complex or a gene splicing
     complex. The present sequence represents a putative novel human
 CC
     spliceosome associated protein (SAP) isolated by the methods of the
 CC
     invention. Note: The prsent sequence is not shown in the specification
 CC
     but was obtained from Genbank or Swissprot using the information provided
 CC
 CC
     in table 2 of the specification
 XX
 SO
     Sequence 448 AA;
                       99.5%; Score 2010; DB 6; Length 448;
  Query Match
  Best Local Similarity
                             Pred. No. 8.9e-203;
                       99.5%;
  Matches 370; Conservative
                             0; Mismatches
                                            2: Indels
                                                           Gaps
                                                                  0;
           1 MDVLSPLSFIKVSHVRMQGILLLVFAKYQHLPYIQILSTKSTPTGLFGYWGNKGGVNICL 60
 Qу
             77 MDVLSPLSFIKVSHVRMOGILLLVFAKYOHLPYIOILSTKSTPTGLFGYWGNKGGVNICL 136
 Db
          61 KLYGYYVSIINCHLPPHISNNYQRLEHFDRILEMQNCEGRDIPNILDHDLIIWFGDMNFR 120
 Qу
             137 KLYGYYVSIINCHLPPHISNNYQRLEHFDRILEMQNCEGRDIPNILDHDLIIWFGDMNFR 196
 Db
         121 IEDFGLHFVRESIKNRCYGGLWEKDQLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDY 130
 QУ
             197 IEDFGLHFVRESIKNRCYGGLWEKDOLSIAKKHDPLLREFOEGRLLFPPTYKFDRNSNDY 256
 Db
         181 DTSEKKRKPAWTDRILWRLKROPCAGPDTPIPPASHFSLSLRGYSSHMTYGISDHKPVSG 240
 QУ
             257 DTSEKKRKPAWTDRILWRLKROPCAGPDTPIPPASHFSLSLRGYSSHMTYGISDHKPVSG 316
 Db
·Qy
         241 TFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVNDY 300
             317 TFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVNDY 376
ď.
         301 VSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFQIPPGSL 360
 Qу
             Db
         377 VSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYRNSLRSVVGIRRPFQIPPGSL 436
         361 REDPLGEAQPQI 372
 QУ
             437 REDPLGEAOPOI 448
· Db
 RESULT 8
 ADN61442
 ID
     ADN61442 standard; protein; 240 AA.
 XX
 AC
     ADN61442;
 XX
 DT
     12-AUG-2004 (first entry)
 XX
 DE
     Human KPP-8 protein SEQ ID NO:8.
 XX
 KW
     human; kinase; phosphatase; enzyme; KPP; cytostatic;
 KW
     antiarteriosclerotic; anticonvulsant; nootropic; neuroprotective;
```

abnormal RNP complexes, (M2) is useful for treating a subject having a

CC

```
KW
     cerebroprotective; anti-HIV; antiallergic; antiinflammatory;
KW
     thyromimetic; gene therapy; cell proliferative disorder; cancer;
KW
     atherosclerosis; neurological disorder; epilepsy; Huntington's disease;
     stroke; immune disorder; inflammatory disorder; AIDS; allergy;
KW
     developmental disorder; Hypothyroidism; Cushing's syndrome; infection.
KW
XX
os
     Homo sapiens.
XX
PN
     WO2004042022-A2.
XX
PD
     21-MAY-2004.
XX
     30-OCT-2003; 2003WO-US034809.
PF
XX
PR
     01-NOV-2002; 2002US-0423226P.
     15-NOV-2002; 2002US-0426713P.
PR
     26-NOV-2002; 2002US-0429766P.
PR
     11-FEB-2003; 2003US-0447043P.
PR
XX
P.A
     (INCY-) INCYTE CORP.
XX
PΙ
     Hafalia AJA, Lee S, Murage J, Swarnakar A, Chawla NK, Khare R;
     Elliott VS, Tran UK, Ramkumar J, Gururajan R, Baughn MR, Gietzen KJ;
PΙ
ΡI
     Yand YG, Chien D, Wang JT, Favero KD, Becha SD, Richardson TW;
PI
     Jin P, Hawkins PR, Yue H, Lee EA, Marquis JP;
XX
DR
     WPI; 2004-390608/36.
     N-PSDB: ADN61498.
DR.
XX
PΤ
     New human kinases and phosphatases (KPP), useful for diagnosing, treating
     and preventing diseases or conditions associated with the aberrant KPP
PT
PT
     expression e.g. cancer, AIDS, epilepsy, or infections.
XX
PS
     Claim 1; SEQ ID NO 8; 320pp; English.
XX
CC
     The present sequence represents a human kinase and phosphatase protein
CC
     designated KPP-8. Human KPP sequences have cytostatic,
CC
     antiarteriosclerotic, anticonvulsant, nootropic, neuroprotective,
CC
     cerebroprotective, anti-HIV, antiallergic, antiinflammatory and
CC
     thyromimetic activities, and can be used in gene therapy. The human KPP
CC
     polypeptides and polynucleotides of the invention are useful in
CC
     diagnosing, treating and preventing diseases or conditions associated
CC
     with the decreased expression or overexpression of KPP, such as cell
CC
     proliferative (e.g. cancer, atherosclerosis), neurological (e.g.
CC
     epilepsy, Huntington's disease, stroke), immune/inflammatory (e.g. AIDS,
CC
     allergies) and developmental (e.g. Hypothyroidism, Cushing's syndrome)
CC
     disorders, or infections. They are also useful in assessing the effects
CC
     of exogenous compounds on the expression of nucleic acid and amino acid
CC
     sequences of KPP. The KPP sequences or their fragments are useful in
CC
     screening compounds for effectiveness as agonist or antagonist of the
CC
     polypeptides, or in altering the expression of the target polynucleotide
CC
     and compounds that specifically bind to or modulate the activity of the
CC
     polypeptide.
XX
SQ
     Sequence 240 AA;
```

```
Best Local Similarity 97.0%; Pred. No. 3.6e-99;
 Matches 191; Conservative
                             1; Mismatches
                                            5: Indels
                                                                     0;
         176 NSNDYDTSEKKRKPAWTDRILWRLKRQPCAGPDTPIPPASHFSLSLRGYSSHMTYGISDH 235
Qу
            Db
         44 NLDIYVIGEKKRKPAWTDRILWRLKROPCAGPDTPIPPASHFSLSLRGYSSHMTYGISDH 103
         236 KPVSGTFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLR 295
QУ
            104 KPVSGTFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLR 163
ďď
         296 DVNDYVSYAWVGDSKVSCSDNLNOVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFOI 355
Qу
            164 DVNDYVSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFQI 223
Db
         356 PPGSLREDPLGEAQPQI 372
Qу
            1111111111111
         224 PPGSLREDPLGEAQPQI 240
Db
RESULT 9
AAU98903
    AAU98903 standard; protein; 1001 AA.
XX
AC
    AAU98903;
XX
DT
    22-AUG-2002 (first entry)
XX
DE
    Rat inositol polyphosphate 5-phosphatase.
XX
KW
    Human; inositol polyphosphate 5-phosphatase; asthma; diabetes;
KW
    chronic obstructive pulmonary disease; cancer; enzyme.
XX
OS
    Rattus norvegicus.
XX
PN
    WO200236755-A2.
XX
PD
    10-MAY-2002.
XX
    29-OCT-2001; 2001WO-EP012496.
PF
XX
PR
    30-OCT-2000; 2000US-0243745P.
    26-DEC-2000: 2000US-0257302P.
PR
    27-AUC-2001: 2001US-0314660P.
PR
XX
PA
    (FARB.) BAYER AG.
XX
ΡI
    Kossida S;
XX
DR
    WPI; 2002-490008/52.
XX
PT
    Now human inositol polyphosphate 5-phosphate polypeptide, useful for
PΤ
    treating chronic obstructive pulmonary disease, asthma, diabetes or
PT
    cancer.
XX
PS
    Disclosure; Fig 3; 135pp; English.
XX
```

```
The invention relates to novel purified human inositol polyphosphate 5-
CC
CC
    phosphatase polypeptide (I) and the polynucleotide (II) encoding it. (I)
    and (II) are useful for screening for agents which decrease the activity
CC
    of human inositol polyphosphate 5-phosphatase polypeptide. (I) or (II)
CC
    are also useful for screening for agents which modulate the activity of
CC
    human inositol polyphosphate 5-phosphatase. A reagent that modulates (I)
CC
    or (II) is useful for treating a human inositol polyphosphate 5-
CC
    phosphatase dysfunction related disease, such as chronic obstructive
CC
    pulmonary disease, asthma, diabetes or cancer. The present sequence
CC.
    represents rat inositol polyphosphate 5- phosphatase
CC
XX
    Sequence 1001 AA;
SO
 Query Match
                        43.2%; Score 872.5; DB 5;
                                                  Length 1001;
 Best Local Similarity
                        47.5%; Pred. No. 4.2e-82;
 Matches 169; Conservative 56; Mismatches 126; Indels
                                                            5;
                                                               Gaps
                                                                       4;
           1 MDVLSPLSFIKVSHVRMQGILLLVFAKYQHLPYIQILSTKSTPTGLFGYWGNKGGVNICL 60
QУ
             483 MDALGPFNFVLVSTVRMQGVILLLFAKYYHLPFLRDVQTDCTRTGLGGYWGNKGGVSVRL 542
Db
Qy
          61 KLYGYYVSIINCHLPPHISNNYQRLEHFDRILEMQNCEGRDIPNILDHDLIIWFGDMNFR 120
               :|: : :||||| |:
                                543 AAFGHMLCFLNCHLPAHMDKAEQRKDNFQTILSLQQFQGPGAHGILDHDLVFWFGDLNFR 602
Db
         121 IEDFGLHFVRESIKNRCYGGLWEKDOLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDY 180
QУ
             11 : | | | | : | :
                               603 IESYDLHFVKFAIDSNQLHQLWEKDQLNMAKNTWPILKGFQEGPLNFAPTFKFDVGTNKY 662
Db
         131 DTSEKKRKPAWTDRILWRLKRQPCAGPDTPIPPASH-FSLSLRGYSSHMTYGISDHKPVS 239
Çу
                                               663 DTSAKKRKPAWTDRILWKVK-APSGGP-SPSGRESHRLQVTQHSYRSHMEYTVSDHKPVA 720
áŒ
         240 GTFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVND 299
QУ
                         721 ARFLLOFAFRDDVPLVRLEVADEWARPEOAVVRYRVETVFARSSWDWIGLYRVGFRHCKD 780
ďď
         300 YVSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFQI 355
Qу
                                     ::|
                                           :|:| |||:
                                                       ::|:: |||
                      : |
                            |\cdot|
         781 YVAYVWAKHEEV--DGNIYQVTFSEESLPKGHGDFILGYYSHHHSILIGVTEPFOI 834
Db
RESULT 10
ADR08504
ID
    ADR08504 standard; protein; 638 AA.
XX
AC
    ADR08504;
XX
DT
    04-NOV-2004 (first entry)
XX
    Human protein useful for treating neurological disease Seq 2010.
DΕ
XX
KW
    human; oligo-capping method; diagnostic marker; gene therapy;
    osteoporosis; neurological disease; Alzheimer's disease;
KW
    Parkinson's disease; dementia; short memory; cancer;
KW
KW
    sense or motor function; emotional reaction; fear response; panic;
KW
    osteopathic; neuroprotective; nootropic; antiparkinsonian; cytostatic;
```

```
KW
     tranquiliser.
XX
os
    Homo sapiens.
XX
PN
    EP1447413-A2.
XX
PD
    18-AUG-2004.
XX
    12-FEB-2004; 2004EP-00003145.
ΡF
XX
     14-FEB-2003; 2003JP-00102207.
PR
PR
     09-MAY-2003; 2003JP-00131452.
XX
PA
     (REAS-) RES ASSOC BIOTECHNOLOGY.
XX
PΙ
     Isogai T, Yamamoto J, Nishikawa T, Isono Y, Sugiyama T, Otsuki T;
ΡI
    Wakamatsu A, Ishii S, Nagai K, Irie R;
XX
DR
    WPI; 2004-583265/57.
DR
    N-PSDB; ADR06548.
XX
    New 1995 cDNA, useful for treating osteoporosis, neurological diseases,
PT
    Alzheimer's diseases, Parkinson's diseases, dementia and various cancers.
PT
XX
PS
    Claim 1; SEQ ID NO 2010; 2686pp; English.
XX
CC
    This invention relates to novel, isolated full length human cDNA
CC
    molecules and the encoded proteins thereof. Specifically, it refers to
CC
     cDNA clones obtained by an oligo-capping method, where none of these
CC
    clones are identical to any known human mRNAs. The present invention
CC
    describes an immunoassay to identify agonists and antagonists, as well as
CC
     antibodies, antisense molecules and siRNAs that can all be used to bind
CC
     to and modulate expression of the cDNA molecules. As such, these
CC
    molecules are useful for diagnostic markers or therapeutic targets for
CC
     the various diseases or morbid states. In particular, they are useful in
CC
     gene therapy for treating osteoporosis, neurological disease, Alzheimer's
CC
     disease, Parkinson's disease, dementia, short memory and various cancers,
CC
     as well as for maintaining equilibrium of sense or motor function, and
CC
     for treating emotional reaction, fear response and panic. Accordingly,
     they exhibit osteopathic, neuroprotective, nootropic, antiparkinsonian,
CC
     cytostatic and tranquiliser activities. This polypeptide is a protein
CC
CC
     encoded by a full length human cDNA sequence of the invention. NOTE: This
CC
     sequence is not given in the sequence listing of the specification but
CC
     can be obtained on CD-ROM from the European Patent Office, Vienna Sub-
CC
    office.
XX
SO
     Sequence 638 AA;
  Query Match
                         43.1%; Score 870.5; DB 8; Length 638;
 Best Local Similarity
                         47.8%; Pred. No. 3.4e-82;
 Matches 170; Conservative 54; Mismatches 127;
                                                                5;
                                                                   Gaps
                                                                            4;
                                                      Indels
            1 MDVLSPLSFIKVSHVRMQGILLLVFAKYQHLPYIQILSTKSTPTGLFGYWGNKGGVNICL 60
Qy
              120 MDALGPFNFVLVSSVRMQGVILLLFAKYYHLPFLRDVQTDCTRTGLGGYWGNKGGVSVRL 179
Db
          61 KLYGYYVSIINCHLPPHISNNYQRLEHFDRILEMQNCEGRDIPNILDHDLIIWFGDMNFR 120
Qy .
```

```
180 AAFGHMLCFLNCHLPAHMDKAEQRKDNFQTILSLQQFQGPGAQGILDHDLVFWFGDLNFR 239
Db
         121 IEDFGLHFVRESIKNRCYGGLWEKDQLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDY 180
Qу
                              || : ||||: :| :
         240 IESYDLHFVKFAIDSDQLHQLWEKDQLNMAKNTWPILKGFQEGPLNFAPTFKFDAGTNKY 299
Db
         181 DTSEKKRKPAWTDRILWRLKROPCAGPDTPIPPASH-FSLSLRGYSSHMTYGISDHKPVS 239
Oy
            300 DTSAKKRKPAWTDRILWKVK-APGGGP-SPSGRKSHRLQVTQHSYRSHMEYTVSDHKPVA 357
Db
         240 GTFDLELKPLVSAPLTVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVND 299
Qу
                              :| | : | | | | | | | | | | | |
                        358 AOFLLOFAFRDDMPLVRLEVADEWVRPEQAVVRYRMETVFARSSWDWIGLYRVGFRHCKD 417
Db
       300 YVSYAWVGDSKVSCSDNLNOVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFOI 355
Qу
                        ::|
                                         :|:| |||::
                                                     ::||: ||||
             418 YVAYVWAKHEDV--DGNTYQVTFSEESLPKGHGDFILGYYSHNHSILIGITEPFQI 471
Db
RESULT 11
ADB65114
    ADE65114 standard; protein; 639 AA.
XX
AC
    ADB65114;
XX
DT
    04-DEC-2003 (first entry)
XX
    Human protein encoded by clone SPLEN20024930.
DE
ХX
    Human; pharmaceutical; diagnostic; gene therapy; tissue regeneration;
KW
KW.
    cell regeneration; membrane protein; signal transduction-related protein;
KW
    transcription-related protein; osteoporosis; neurological disease;
    cancer; tumour.
KW
XX
    Homo sapiens.
os
XX
PN
    EP1308459-A2.
XX
PD
    07-MAY-2003.
XΧ
    28-MAR-2002; 2002EP-00007401.
PF
XX
PR
    05-NOV-2001; 2001JP-00379298.
    25-JAN-2002; 2002US-00350978.
PR
XX
PA
     (HELI-) HELIX RES INST.
     (REAS-) RES ASSOC BIOTECHNOLOGY.
PΑ
XX
PΙ
    Isoqai T, Sugiyama T, Otsuki T, Wakamatsu A, Sato H, Ishii S;
    Yamamoto J, Isono Y, Hio Y, Otsuka K, Nagai K, Irie R, Tamechika I;
PΙ
    Seki N, Yoshikawa T, Otsuka M, Nagahari K, Masuho Y;
PΙ
XX
    WPI; 2003-450961/43.
DR
    N-PSDB; ADB63144.
DR
XX
    New polynucleotides and polypeptides, useful for developing a diagnostic
PΤ
```

marker or medicines for regulation of their expression and activity, or as targets of gene therapy.

PT XX PS

PT

Claim 1; Page; 222pp; English.

XX CC

The invention discloses a polynucleotide comprising a sequence selected from 1970 fully defined nucleotide sequences which encode novel polypeptides. Also claimed is a polypeptide encoded by the polynucleotide or its partial peptide, an antibody binding to the polypeptide or peptide of the polynucleotide, immunologically assaying the polypeptide or peptide of the polynucleotide by contacting the polypeptide or peptide with the antibody of the encoded protein, and observing the binding between the two, a transformant carrying the polynucleotide in an expressible manner and an antisense polynucleotide. The oligonucleotide is useful as a primer for synthesising the polynucleotide, or as a probe for detecting the polynucleotide. The polynucleotides and encoded proteins are useful as pharmaceutical agents and many disease-related genes may be included in them, for developing a diagnostic marker or medicines for regulation of their expression and activity, or as targets of gene therapy. The genes are involved in tissue and/or cell regeneration. Membrane proteins, signal transduction-related proteins, transcription-related proteins, disease-related proteins and genes encoding them can be used as indicators for diseases (e.g. osteoporosis, neurological diseases, cancer, tumours. The cDNA may be used to regulate the activity or expression of the encoded protein to treat diseases. The sequence presented is a protein of the invention. Note: Some of the sequence data for this patent is not represented in the printed specification, but is based on sequence information supplied by the European Patent Office.

43.0%; Score 868.5; DB 7; Length 639;

CC XX SQ

Sequence 639 AA;

Query Match

```
Best Local Similarity
                     47.8%; Pred. No. 5.6e-82;
 Matches 170; Conservative 54; Mismatches 127; Indels
                                                      5; Gaps
          1 MDVLSPLSFIKVSHVRMQGILLLVFAKYQHLPYIQILSTKSTPTGLFGYWGNKGGVNICL 60
Qу
           121 MDALGPFNFVLVSSVRMQGVILLLFAKYYHLPFLRDVQTDCTRTGLGGYWGNKGGVSVRL 180
Db
         61 KLYGYYVSIINCHLPPHISNNYORLEHFDRILEMONCEGRDIPNILDHDLIIWFGDMNFR 120
QУ
                             :|: : :|||||
        181 AAFGHMLCFLNCHLPAHMDKAEORKDNFOTILSLOOFOGPGAOGILDHDLVFWFGDLNFR 240
Db
        121 IEDFGLHFVRESIKNRCYGGLWEKDOLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDY 180
QУ
                            || : ||||: :| :
        241 IESYDLHFVKFAIDSDOLHOLWEKDOLNMAKNTWPILKGFQEGPLNFAPTFKFDVGTNKY 300
Db
        181 DTSEKKRKPAWTDRILWRLKROPCAGPDTPIPPASH-FSLSLRGYSSHMTYGISDHKPVS 239
Qy
           301 DTSAKKRKPAWTDRILWKVK-APGGGP-SPSGRKSHRLQVTQHSYRSHMEYTVSDHKPVA 358
Db
        240 GTFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVND 299
Qу
                      |\cdot|: |
             | |:
                             : | | : | | | | | | | | | | | | | |
        359 AQFLLQFAFRDDMPLVRLEVADEWVRPEQAVVRYRMETVFARSSWDWIGLYRVGFRHCKD 418
Db
        300 YVSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFOI 355
QУ
```

```
RESULT 12
ADM04740
     ADM04740 standard; protein; 639 AA.
ID
XX
AC
     ADM04740;
XX
DT
     20-MAY-2004 (first entry)
XX
DE
     Human protein of the invention SEQ ID NO:3425.
XX
KW
     human; gene therapy; diagnostic marker; pharmaceutical.
XX
os
     Homo sapiens.
XX
PN
     EP1347046-A1.
XX
PD
     24-SEP-2003.
XX
PF
     12-APR-2002; 2002EP-00008400.
XX
PR
     22-MAR-2002; 2002JP-00137785.
XX
     (REAS-) RES ASSOC BIOTECHNOLOGY.
PΑ
XX
PΙ
     Isogai T, Sugiyama T, Otsuki T, Wakamatsu A, Sato H, Ishii S;
PΙ
     Yamamoto J, Isono Y, Hio Y, Otsuka K, Nagai K, Irie R, Tamechika I;
     Seki N, Yoshikawa T, Otsuka M, Nagahari K, Masuho Y;
PΙ
ΧХ
DR
     WPI; 2003-723553/69.
DR
     N-PSDB: ADM02297.
XX
PT
    New polynucleotides and polypeptides are useful in gene therapy, for
PT
     developing a diagnostic marker or medicines for regulating their
PT
     expression and activity, or as a target of gene therapy.
XX
PS
     Claim 1; SEQ ID NO 3425; 305pp; English.
XX
CC
     The invention relates to a novel human polynucleotide and the encoded
CC
     polypeptide. A polynucleotide of the invention may have a use in gene
CC
     therapy. An oligonucleotide of the invention ADM06202-ADM06773 is useful
CC
     as a primer for synthesizing the polynucleotide or as a probe for
CC
     detecting the polynucleotide. The polynucleotides ADM01316-ADM03758 are
    useful in gene therapy, for developing a diagnostic marker or medicines
CC
CC
     for regulating their expression and activity, or as a target of gene
CC
     therapy. The proteins ADM03759-ADM06201 encoded by the polynucleotides
CC
     are useful as pharmaceutical agents. The present sequence represents a
CC.
     protein sequence of the invention.
XX
SQ
    Sequence 639 AA;
 Query Macch
                          43.0%; Score 868.5; DB 7; Length 639;
 Best Local Similarity
                          47.8%; Pred. No. 5.6e-82;
 Matches 170; Conservative 54; Mismatches 127; Indels
```

```
1 MDVLSPLSFIKVSHVRMOGILLLVFAKYOHLPYIOILSTKSTPTGLFGYWGNKGGVNICL 60
Qу
            121 MDALGPFNFVLVSSVRMQGVILLLFAKYYHLPFLRDVQTDCTRTGLGGYWGNKGGVSVRL 180
Db
         61 KLYGYYVSIINCHLPPHISNNYORLEHFDRILEMONCEGRDIPNILDHDLIIWFGDMNFR 120
QУ
              :|: : :||||| |: || || ::| || :| :|
                                                   181 AAFGHMLCFLNCHLPAHMDKAEQRKDNFQTILSLQQFQGPGAQGILDHDLVFWFGDLNFR 240
Db
        121 IEDFGLHFVRESIKNRCYGGLWEKDQLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDY 180
Qу
                             || : ||||: :| :
        241 IESYDLHFVKFAIDSDOLHOLWEKDOLNMAKNTWPILKGFQEGPLNFAPTFKFDVGTNKY 300
Db
QУ
        181 DTSEKKRKPAWTDRILWRLKROPCAGPDTPIPPASH-FSLSLRGYSSHMTYGISDHKPVS 239
            ::
                                                   | | | | | | : | | | | | | :
        301 DTSAKKRKPAWTDRILWKVK-APGGGP-SPSGRKSHRLQVTQHSYRSHMEYTVSDHKPVA 358
Db
Qу
        240 GTFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVND 299
              1 :
                       359 AQFLLQFAFRDDMPLVRLEVADEWVRPEQAVVRYRMETVFARSSWDWIGLYRVGFRHCKD 418
Db
Qy
        300 YVSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFQI 355
            419 YVAYVWAKHEDV--DGNTYOVTFSEESLPKGHGDFILGYYSHNHSILIGITEPFOI 472
Db
RESULT 13
AAU98904
ID
    AAU98904 standard; protein; 1006 AA.
XX
AC
    AAU98904;
XX
DT
    22-AUG-2002 (first entry)
XX
DE
    Human inositol polyphosphate 5-phosphatase #2.
XX
KW
    Human; inositol polyphosphate 5-phosphatase; asthma; diabetes;
ΚW
    chronic obstructive pulmonary disease; cancer; enzyme.
XX
os
    Homo sapiens.
XX
ĎΝ
    WO200236755-A2.
XX
PD
    10-MAY-2002.
XX
PF
    29-OCT-2001; 2001WO-EP012496.
XX
PR
    30-OCT-2000; 2000US-0243745P.
PR
    26-DEC-2000; 2000US-0257302P.
PR
    27-AUG-2001; 2001US-0314660P.
XX
PA
    (FARE ) BAYER AG.
XX
ΡI
    Kossida S;
XX
    WPI; 2002-490008/52.
DR
DR
    N-PSDB; ABK86166.
```

```
PT
    New human inositol polyphosphate 5-phosphate polypeptide, useful for
PT
    treating chronic obstructive pulmonary disease, asthma, diabetes or
PT
    cancer.
XX
PS
    Claim 25; Fig 12; 135pp; English.
XX
CC
    The invention relates to novel purified human inositol polyphosphate 5-
    phosphatase polypeptide (I) and the polynucleotide (II) encoding it. (I)
CC
    and (II) are useful for screening for agents which decrease the activity
CC
    of human inositol polyphosphate 5-phosphatase polypeptide. (I) or (II)
CC
    are also useful for screening for agents which modulate the activity of
CC
    human inositol polyphosphate 5-phosphatase. A reagent that modulates (I)
CC
CC
    or (II) is useful for treating a human inositol polyphosphate 5-
CC
    phosphatase dysfunction related disease, such as chronic obstructive
CC
    pulmonary disease, asthma, diabetes or cancer. The present sequence
CC
    represents human inositol polyphosphate 5- phosphatase
XX
SO
    Sequence 1006 AA;
 Query Match
                        43.0%; Score 868.5; DB 5; Length 1006;
 Best Local Similarity 47.8%; Pred. No. 1.1e-81;
 Matches 170; Conservative 54; Mismatches 127;
                                                   Indels
                                                             5;
                                                                Gaps
                                                                        4;
           1 MDVLSPLSFIKVSHVRMOGILLLVFAKYOHLPYIOILSTKSTPTGLFGYWGNKGGVNICL 60
Qy
             Db
         488 MDALGPFNFVLVSSVRMOGVILLLFAKYYHLPFLRDVOTDCTRTGLGGYWGNKGGVSVRL 547 ·
          61 KLYGYYVSIINCHLPPHISNNYORLEHFDRILEMONCEGRDIPNILDHDLIIWFGDMNFR 120
QУ
                                 || ::| || :| :|
               :|: : :||||| |:
                                                      548 AAFGHMLCFLNCHLPAHMDKAEORKDNFQTILSLQOFQGPGAQGILDHDLVFWFGDLNFR 607
Db
         121 IEDFGLHFVRESIKNRCYGGLWEKDOLSIAKKHDPLLREFOEGRLLFPPTYKFDRNSNDY 180
Qу
             || : ||||: :| :
                                |||||||
                                            608 IESYDLHFVKFAIDSDQLHQLWEKDQLNMAKNTWPILKGFQEGPLNFAPTFKFDVGTNKY 667
Db
         181 DTSEKKRKPAWTDRILWRLKRQPCAGPDTPIPPASH-FSLSLRGYSSHMTYGISDHKPVS 239
2у
             ::
         668 DTSAKKRKPAWTDRILWKVK-APGGGP-SPSGRKSHRLQVTQHSYRSHMEYTVSDHKPVA 725
Db
         240 GTFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVND 299
QУ
                                         :| | : | | | | | | | | | | | |
                               | | |
         726 AOFLLOFAFRDDMPLVRLEVADEWVRPEOAVVRYRMETVFARSSWDWIGLYRVGFRHCKD 785
Db
         300 YVSYAWVGDSKVSCSDNLNOVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFQI 355
Qу
                           ::|
                                            :|:| |;|::
         786 YVAYVWAKHEDV--DGNTYOVTFSEESLPKGHGDFILGYYSHNHSILIGITEPFOI 839
Db
RESULT 14
ABU54583
    ABU54583 standard; protein; 1006 AA.
TD
XX
AC
    ABU54583;
XX
DT
    03-JUN-2003 (first entry)
XX
```

XX

```
DE
     Human NOVX polypeptide #42.
XX
KW
     Human; NOVX; metabolic disorder; cardiomyopathy; diabetes; ASD;
KW
     hypertension; congenital heart defect; aortic stenosis; valve disease;
     atrial septal defect; atrioventricular canal defect; ductus arteriosus;
KW
KW
     pulmonary stenosis; subaortic stenosis; ventricular septal defect; VSD;
KW
     tuberous sclerosis; scleroderma; atherosclerosis; infectious disease;
     obesity; anorexia; neurodegenerative disorder; Alzheimer's disease;
KW
KW
     Parkinson's disease; immune disorder; haematopoietic disorder;
KW
     haemophilia; hypercoagulation; Crohn's disease; cancer.
XX
OS
     Homo sapiens.
XX
PN
     WO200281498-A2.
XX
PD
     17-OCT-2002.
XX
PF
     03-APR-2002; 2002WO-US010780.
XX
PR
     03-APR-2001; 2001US-0281086P.
PR
     03-APR-2001; 2001US-0281136P.
PR
     05-APR-2001; 2001US-0281863P.
     05-APR-2001; 2001US-0281906P.
PR
     06-APR-2001; 2001US-0282020P.
PR
     10-APR-2001; 2001US-0282930P.
₽R
PR
     10-APR-2001; 2001US-0282934P.
PR
     12-APR-2001; 2001US-0283512P.
PR
     13-APR-2001; 2001US-0283710P.
PR
     17-APR-2001; 2001US-0284234P.
PR
     19-APR-2001; 2001US-0285325P.
     20-APR-2001; 2001US-0285381P.
PR
PR
     20-APR-2001; 2001US-0285609P.
PR
     23-APR-2001; 2001US-0285748P.
PR
     23-APR-2001; 2001US-0285890P.
PR
     24-APR-2001; 2001US-0286068P.
PR
     25-APR-2001; 2001US-0286292P.
PR-
     27-APR-2001; 2001US-0287213P.
PR
     02-MAY-2001; 2001US-0288257P.
PR
     29-MAY-2001; 2001US-0294164P.
PR
     30-MAY-2001; 2001US-0294484P.
PR
     18-JUN-2001; 2001US-0298952P.
PR
     19-JUN-2001; 2001US-0299237P.
PR
     19-JUN-2001; 2001US-0299276P.
PR
     12-SEP-2001; 2001US-0318750P.
     25-SEP-2001; 2001US-0324800P.
PR
     25-SEP-2001; 2001US-0324802P.
PR
PR
     27-SEP-2001; 2001US-0325684P.
PR
     17-OCT-2001; 2001US-0330143P.
PR
     14-NOV-2001; 2001US-0332131P.
PR.
     14-NOV-2001; 2001US-0332240P.
     14-NOV-2001; 2001US-0332779P.
PR
     21-NOV-2001; 2001US-0332115P.
₽R
PR
     04-DEC-2001; 2001US-0337621P.
PR
     03-JAN-2002; 2002US-0345783P.
PR
    · 16-JAN-2002; 2002US-0350251P.
PR
     02-APR-2002; 2002US-00114270.
XX
```

```
PΑ
     (CURA-) CURAGEN CORP.
XX
    Guo X, Kekuda R, Miller CE, Malyankar UM, Spytek KA;
PΙ
    Patturajan M, Liu X, Gusev VY, Li L, Vernet CAM, Zerhusen BD;
ΡI
    Gorman L, Shenoy SG, Pena CEA, Smithson G, Burgess CE, Gerlach V;
PΙ
    Padigaru M, Shimkets RA, Gangolli EA, Taupier RJ, Casman SJ, Ji W;
PΙ
    Anderson DW, Leite MW, Rastelli L, Edinger SR, Stone DJ;
PI
    Macdougall JR, Rothenberg ME, Mazur A, Millet I, Peyman JA;
PΙ
PΙ
    Ellerman K;
XX
DR
    WPI; 2003-046858/04.
DR
    N-PSDB; ABX72211.
XX
PT
    New isolated NOVX polypeptide useful for treating atherosclerosis,
PT
    metabolic disorders, diabetes, obesity, infectious disease, anorexia,
    neurodegenerative disorders, Alzheimer's disease and cancer.
PT
XX
PS
    Claim 1; Page 179-180; 666pp; English.
XX
    The invention relates to human polypeptides, termed NOVX, and the
CC
CC
    polynucleotides encoding them. The polypeptides and polynucleotides are
    useful for diagnosing disease, and screening for potential therapeutic
CC
CC
    agents. The sequences are useful for treating metabolic disorders,
    cardiomyopathy, diabetes, hypertension, congenital heart defects, aortic
CC
    stenosis, atrial septal defect (ASD), atrioventricular canal defect,
CC
    ductus arteriosus, pulmonary stenosis, subaortic stenosis, ventricular
CC
    septal defect (VSD), valve diseases, tuberous sclerosis, scleroderma,
CC
    atherosclerosis, obesity, infectious disease, anorexia, neurodegenerative
CC
    disorders, Alzheimer's disease, Parkinson's disease, immune disorders,
CC
    haematopoietic disorders, haemophilia, hypercoagulation, Crohn's disease
CC
CC
    and cancer. Sequences ABU54542-ABU54647 represent human NOVX polypeptides
    of the invention
CC
XX
SO
    Sequence 1006 AA;
                        43.0%; Score 868.5; DB 6; Length 1006;
  Query Match
                        47.8%; Pred. No. 1.1e-81;
  Best Local Similarity
  Matches 170; Conservative 54; Mismatches 127; Indels
                                                            5; Gaps
                                                                       4;
           1 MDVLSPLSFIKVSHVRMQGILLLVFAKYQHLPYIQILSTKSTPTGLFGYWGNKGGVNICL 60
Qу
             438 MDALGPFNFVLVSSVRMQGVILLLFAKYYHLPFLRDVQTDCTRTGLGGYWGNKGGVSVRL 547
Db
          61 KLYGYYVSIINCHLPPHISNNYQRLEHFDRILEMQNCEGRDIPNILDHDLIIWFGDMNFR 120
QУ
               :|: : :||||| |:
                                 |||||||
         548 AAFGHMLCFLNCHLPAHMDKAEQRKDNFQTILSLQQFQGPGAQGILDHDLVFWFGDLNFR 607
Dl_2
         121 IEDFGLHFVRESIKNRCYGGLWEKDQLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDY 180
Qу
                               || : ||||: :| :
         603 IESYDLHFVKFAIDSDQLHQLWEKDQLNMAKNTWPILKGFQEGPLNFAPTFKFDVGTNKY 667
Db
         181 DTSEKKRKPAWTDRILWRLKROPCAGPDTPIPPASH-FSLSLRGYSSHMTYGISDHKPVS 239
QУ
             ::
         668 DTSAKKRKPAWTDRILWKVK-APGGGP-SPSGRKSHRLQVTQHSYRSHMEYTVSDHKPVA 725
Db
         240 GTFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVND 299
Qу
```

```
Db
          726 AQFLLQFAFRDDMPLVRLEVADEWVRPEQAVVRYRMETVFARSSWDWIGLYRVGFRHCKD 785
          300 YVSYAWVGDSKVSCSDNLNOVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFOI 355
Qу
                              1 11
                                               : | : | | | | | : : |
                                       ::|
                                                            ::||: ||||
          786 YVAYVWAKHEDV--DGNTYQVTFSEESLPKGHGDFILGYYSHNHSILIGITEPFQI 839
Db
RESULT 15
AAU98902
     AAU98902 standard; protein; 751 AA.
XX
AC
     AAU98902;
ХX
DT
     22-AUG-2002 (first entry)
XX
DE
     Human inositol polyphosphate 5-phosphatase #1.
XX
     Human; inositol polyphosphate 5-phosphatase; asthma; diabetes;
KW
KW
     chronic obstructive pulmonary disease; cancer; enzyme.
XX
OS
     Homo sapiens.
XX
PN
     WO200236755-A2.
XX
PD
     10-MAY-2002.
XX
PF
     29-OCT-2001; 2001WO-EP012496.
XX
PR
     30-OCT-2000; 2000US-0243745P.
PR
     26-DEC-2000; 2000US-0257302P.
     27-AUG-2001; 2001US-0314660P.
PR
XX
PΑ
     (FARB ) BAYER AG.
XX
PΙ
     Kossida S;
XX
DR
     WPI; 2002-490008/52.
DR
     N-PSDB; ABK86158.
XX
PT
     New human inositol polyphosphate 5-phosphate polypeptide, useful for
PT
     treating chronic obstructive pulmonary disease, asthma, diabetes or
PT
     cancer.
XX
PS
     Claim 25; Fig 2; 135pp; English.
XX
CC
     The invention relates to novel purified human inositol polyphosphate 5-
CC
     phosphatase polypeptide (I) and the polynucleotide (II) encoding it. (I)
CC
     and (II) are useful for screening for agents which decrease the activity
CC
     of human inositol polyphosphate 5-phosphatase polypeptide. (I) or (II)
CC
     are also useful for screening for agents which modulate the activity of
CC
     human inositol polyphosphate 5-phosphatase. A reagent that modulates (I)
CC
     or (II) is useful for treating a human inositol polyphosphate 5-
CC
     phosphatase dysfunction related disease, such as chronic obstructive
CC
     pulmonary disease, asthma, diabetes or cancer. The present sequence
CC
     represents a human inositol polyphosphate 5- phosphatase
XX
SO
     Sequence 751 AA;
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41.2%; Score 831.5; DB 5; Length 751; Query Match Best Local Similarity 42.9%; Pred. No. 5.8e-78; Matches 169; Conservative 56; Mismatches 126; Indels 43; Gaps 6; 1 MDVLSPLSFIKVSH------VRMOGILLLVFA 26 QУ || | | : | : | : | | | | | | | : : | | : | | 195 MDALGPFNFVLVTHPSPPGOPETLLNSWLOLYPGSLWGPLGLCGWVSSVRMOGVILLLFA 254 Db 27 KYQHLPYIQILSTKSTPTGLFGYWGNKGGVNICLKLYGYYVSIINCHLPPHISNNYQRLE 86 Qy 255 KYYHLPFLRDVOTDCTRTGLGGYWGNKGGVSVRLAAFGHMLCFLNCHLPAHMDKAEORKD 314 Db 87 HFDRILEMONCEGRDIPNILDHD----LIIWFGDMNFRIEDFGLHFVRESIKNRCYGGLW 142 QУ 315 NFQTILSLQQFQGPGAQGILDHEYGLGLVFWFGDLNFRIESYDLHFVKFAIDSDQLHQLW 374 Db Qу 143 EKDQLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDYDTSEKKRKPAWTDRILWRLKRQ 202 375 EKDQLNMAKNTWPILKGFQEGPLNFAPTFKFDVGTNKYDTSAKKRKPAWTDRILWKVK-A 433 Db ÒУ 203 PCAGPDTPIPPASH-FSLSLRGYSSHMTYGISDHKPVSGTFDLELKPLVSAPLIVLMPED 261 ||: | 434 PGGGP-SPSGRKSHRLOVTOHSYRSHMEYTVSDHKPVAAQFLLQFAFRDDMPLVRLEVAD 492 Db 262 LWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVNDYVSYAWVGDSKVSCSDNLNQVY 321 Qу 493 EWVRPEQAVVRYRMETVFARSSWDWIGLYRVGFRHCKDYVAYVWAKHEDV--DGNTYQVT 550 Db 322 IDISNIPTTEDEFLLCYYSNSLRSVVGISRPFOI 355 Qу Db 551 FSEESLPKGHGDFILGYYSHNHSILIGITEPFQI 584

Search completed: February 10, 2005, 17:49:57 Job time: 170 secs

> GenCore version 5.1.6 Copyright (c) 1993 - 2005 Compugen Ltd.

OM protein - protein search, using sw model

Run on: February 10, 2005, 17:50:07; Search time 43 Seconds

(without alignments)

645.801 Million cell updates/sec

Title: US-09-892-287-1

Perfect score: 2020

Sequence: 1 MDVLSPLSFIKVSHVRMQGI......FQIPPGSLREDPLGEAQFQI 372

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 513545 seqs, 74649064 residues

Total number of hits satisfying chosen parameters: 513545

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : Is:

Issued_Patents_AA:*

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Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

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1	2020	100.0	372	2	US-08-884-681-1	Sequence 1, Appli
2	2020	100.0	372	3	US-09-258-643-1	Sequence 1, Appli
3		88.7	329	2	US-08-884-681-3	Sequence 3, Appli
4	1791	88.7	329	3	US-09-258-643-3	Sequence 3, Appli
5	432.5	21.4	942	2	US-08-884-681-4	Sequence 4, Appli
6	432.5	21.4	942	3	US-08-560-005-6	Sequence 6, Appli
7	432.5	21.4	942	3	US-09-258-643-4	Sequence 4, Appli
8	432.5	21.4	942	3	US-09-418-540-6	Sequence 5, Appli
. 9	432.5	21.4	942	4	US-09-969-528-6	Sequence 6, Appli
10	368.5	18.2	968	3	US-08-560-005-7	Sequence 7, Appli
11	368.5	18.2	968	3	US-09-418-540-7	Sequence 7, Appli
12	368.5	18.2	968	4	US-09-969-528-7	Sequence 7, Appli
13	367.5	18.2	901	2	US-08-884-681-5	Sequence 5, Appli
14	367.5	18.2	901	3	US-09-258-643-5	Sequence 5, Appli
15	367.5	18.2	952	4	US-09-949-016-7283	Sequence 7283, Ap
16	367.5	18.2	952	4	US-09-949-016-7284	Sequence 7284, Ap
17	328	16.2	350	4	US-09-248-796A-18330	Sequence 18330, A
18	328	16.2	1587	4	US-09-949-016-11062	Sequence 11062, A
19	324.5	16.1	818	4	US-09-248-796A-18342	Sequence 18342, A
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ALIGNMENTS

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US-08-834-681-1
; Sequence 1, Application US/08884681
; Patent No. 5955338
  GENERAL INFORMATION:
    APPLICANT: Hillman, Jennifer L.
    APPLICANT: Lal, Preeti
    APPLICANT: Corley, Neil C.
    APPLICANT: Shah, Purvi
    TITLE OF INVENTION: NEW HUMAN PHOSPHATIDYLINOSITOL
    NUMBER OF SEQUENCES: 5
    CORRESPONDENCE ADDRESS:
      ADDRESSEE: Incyte Pharmaceuticals, Inc.
      STREET: 3174 Porter Drive
      CITY: Palo Alto
      STATE: CA
      COUNTRY: USA
      ZIP: 94304
    COMPUTER READABLE FORM:
      MEDIUM TYPE: Diskette
      COMPUTER: IBM Compatible
      OPERATING SYSTEM: DOS
       SOFTWARE: FastSEQ for Windows Version 2.0
     CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/08/884,681
       FILING DATE: Filed Herewith
     PRIOR APPLICATION DATA:
      APPLICATION NUMBER:
       FILING DATE:
    ATTORNEY/AGENT INFORMATION:
      NAME: Billings, Lucy J.
      REGISTRATION NUMBER: 36,749
      REFERENCE/DOCKET NUMBER: PF-0334 US
     TELECOMMUNICATION INFORMATION:
       TELEPHONE: 415-855-0555
       TELEFAX: 415-845-4166
   INFORMATION FOR SEQ ID NO: 1:
     SEQUENCE CHARACTERISTICS:
      LENGTH: 372 amino acids
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STRANDEDNESS: single
     TOPOLOGY: linear
    IMMEDIATE SOURCE:
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     CLONE: 638789
US-08-884-681-1
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US-09-258-643-1
; Sequence 1, Application US/09258643
; Patent No. 6277373
  GENERAL INFORMATION:
    APPLICANT: Hillman, Jennifer L.
    APPLICANT: Lal, Preeti
    APPLICANT: Corley, Neil C.
    APPI:ICANT: Shah, Purvi
    TITLE OF INVENTION: NEW HUMAN PHOSPHATIDYLINOSITOL
    NUMBER OF SEQUENCES: 5
    CORRESPONDENCE ADDRESS:
     ADDRESSEE: Incyte Pharmaceuticals, Inc.
     STREET: 3174 Porter Drive
     CITY: Palo Alto
     STATE: CA
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TYPE: amino acid

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COUNTRY: USA
     ZIP: 94304
    COMPUTER READABLE FORM:
     MEDIUM TYPE: Diskette
     COMPUTER: IBM Compatible
     OPERATING SYSTEM: DOS
     SOFTWARE: FastSEQ for Windows Version 2.0
    CURRENT APPLICATION DATA:
     APPLICATION NUMBER: US/09/258,643
     FILING DATE:
    PRIOR APPLICATION DATA:
     APPLICATION NUMBER: US/08/884,681
     FILING DATE:
    ATTORNEY/AGENT INFORMATION:
     NAME: Billings, Lucy J.
     REGISTRATION NUMBER: 36,749
     REFERENCE/DOCKET NUMBER: PF-0334 US
    TELECOMMUNICATION INFORMATION:
     TELEPHONE: 415-855-0555
     TELEFAX: 415-845-4166
  INFORMATION FOR SEQ ID NO:
    SEQUENCE CHARACTERISTICS:
     LENGTH: 372 amino acids
     TYPE: amino acid
     STRANDEDNESS: single
     TOPOLOGY: linear
    IMMEDIATE SOURCE:
     LIBRARY: BRSTNOT03
     CLONE: 638789
US-09-253-643-1
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RESULT 3
US-08-884-681-3
; Sequence 3, Application US/08884681
; Patent No. 5955338
  GENERAL INFORMATION:
    APPLICANT: Hillman, Jennifer L.
    APPLICANT: Lal, Preeti
    APPLICANT: Corley, Neil C.
    APPLICANT: Shah, Purvi
    TITLE OF INVENTION: NEW HUMAN PHOSPHATIDYLINOSITOL
    NUMBER OF SEQUENCES: 5
    CORRESPONDENCE ADDRESS:
;
      ADDRESSEE: Incyte Pharmaceuticals, Inc.
      STREET: 3174 Porter Drive
     CITY: Palo Alto
     STATE: CA
     COUNTRY: USA
      ZIP: 94304
    COMPUTER READABLE FORM:
      MEDIUM TYPE: Diskette
      COMPUTER: IBM Compatible
      OPERATING SYSTEM: DOS
      SOFTWARE: FastSEO for Windows Version 2.0
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/08/884,681
      FILING DATE: Filed Herewith
    PRIOR APPLICATION DATA:
     APPLICATION NUMBER:
      FILING DATE:
    ATTORNEY/AGENT INFORMATION:
      NAME: Billings, Lucy J.
      REGISTRATION NUMBER: 36,749
      REFERENCE/DOCKET NUMBER: PF-0334 US
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: 415-855-0555
      TELEFAX: 415-845-4166
 INFORMATION FOR SEQ ID NO: 3:
    SEQUENCE CHARACTERISTICS:
      LENGTH: 329 amino acids
      TYPE: amino acid
      STRANDEDNESS: single
      TOPOLOGY: linear
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      LIBRARY: GenBank
      CLONE: 1399101
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; Sequence 3; Application US/09258643
; Patent No. 6277373
  GENERAL INFORMATION:
    APPLICANT: Hillman, Jennifer L.
    APPLICANT: Lal, Preeti
    APPLICANT: Corley, Neil C.
    APPLICANT: Shah, Purvi
    TITLE OF INVENTION: NEW HUMAN PHOSPHATIDYLINOSITOL
    NUMBER OF SEQUENCES: 5
    CORRESPONDENCE ADDRESS:
      ADDRESSEE: Incyte Pharmaceuticals, Inc.
      STREET: 3174 Porter Drive
      CITY: Palo Alto
      STATE: CA
      COUNTRY: USA
      ZIP: 94304
    COMPUTER READABLE FORM:
      MEDIUM TYPE: Diskette
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      OPERATING SYSTEM: DOS
      SOFTWARE: FastSEQ for Windows Version 2.0
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/09/258,643
      FILING DATE:
    PRIOR APPLICATION DATA:
      APPLICATION NUMBER: US/08/884,681
      FILING DATE:
    ATTORNEY/AGENT INFORMATION:
      NAME: Billings, Lucy J.
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REGIS'TRATION NUMBER: 36,749
      REFERENCE/DOCKET NUMBER: PF-0334 US
    TELECOMMUNICATION INFORMATION:
     TELEPHONE: 415-855-0555
      TELEFAX: 415-845-4166
  INFORMATION FOR SEQ ID NO:
    SEQUENCE CHARACTERISTICS:
     LENGTH: 329 amino acids
     TYPE: amino acid
     STRANDEDNESS: single
     TOPOLOGY: linear
    IMMEDIATE SOURCE:
     LIBRARY: GenBank
     CLONE: 1399101
US-09-258-643-3
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                            Score 1791; DB 3; Length 329;
 Best Local Similarity
                     99.4%;
                            Pred. No. 4.7e-184;
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US-08-884-681-4
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 Patent No. 5955338
  GENERAL INFORMATION:
    APPLICANT: Hillman, Jennifer L.
    APPLICANT: Lal, Preeti
    APPLICANT: Corley, Neil C.
    APPLICANT: Shah, Purvi
    TITLE OF INVENTION: NEW HUMAN PHOSPHATIDYLINOSITOL
    NUMBER OF SEQUENCES: 5
    CORRESPONDENCE ADDRESS:
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ADDRESSEE: Incyte Pharmaceuticals, Inc.
     STREET: 3174 Porter Drive
     CITY: Palo Alto
     STATE: CA
     COUNTRY: USA
     ZIP: 94304
    COMPUTER READABLE FORM:
     MEDIUM TYPE: Diskette
     CCMPUTER: IBM Compatible
     OPERATING SYSTEM: DOS
     SOFTWARE: FastSEQ for Windows Version 2.0
   CURRENT APPLICATION DATA:
    APPLICATION NUMBER: US/08/884,681
     FILING DATE: Filed Herewith
    PRIOR APPLICATION DATA:
    APPLICATION NUMBER:
     FILING DATE:
    ATTORNEY/AGENT INFORMATION:
    NAME: Billings, Lucy J.
    REGISTRATION NUMBER: 36,749
    REFERENCE/DOCKET NUMBER: PF-0334 US
    TELECOMMUNICATION INFORMATION:
     TELEPHONE: 415-855-0555
     TELEFAX: 415-845-4166
  INFORMATION FOR SEQ ID NO: 4:
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     STRANDEDNESS: single
     TOPOLOGY: linear
    IMMEDIATE SOURCE:
     LIBRARY: GenBank
     CLCNE: 1019103
US-08-884-681-4
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           DЪ
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Qу
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Qу
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RESULT 6
US-08-560-005-6
; Sequence 6, Application US/08560005
; Patent No. 6001354
  GENERAL INFORMATION:
    APPLICANT: Pot, David A.
    APPLICANT: Williams, Lewis T.
    APPLICANT: Jefferson, Anne Bennett
    APPLICANT: Majerus, Philip W.
    TITLE OF INVENTION: No. 6001354el Grb2 Associating Protein and Nucleic
    TITLE OF INVENTION: Acids Encoding Therefor
    NUMBER OF SEQUENCES: 10
    CORRESPONDENCE ADDRESS:
      ADDRESSEE: Townsend and Townsend and Crew
      STREET: One Market Plaza, Steuart Tower, Suite 2000
      CITY: San Francisco
      STATE: California
      COUNTRY: USA
      ZIP: 94105
    COMPUTER READABLE FORM:
      MEDIUM TYPE: Floppy disk
      COMPUTER: IBM PC compatible
      OPERATING SYSTEM: PC-DOS/MS-DOS
      SOFTWARE: PatentIn Release #1.0, Version #1.25
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/08/560,005
      FILING DATE:
      CLASSIFICATION: 435
    ATTORNEY/AGENT INFORMATION:
      NAME: Dow, Karen B.
      REGISTRATION NUMBER: 29,684
      REFERENCE/DOCKET NUMBER: 2307K-0624000
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: 415-326-2400
      TELEFAX: 415-326-2422
  INFORMATION FOR SEQ ID NO:
    SEQUENCE CHARACTERISTICS:
      LENGTH: 942 amino acids
      TYPE: amino acid
      TOPOLOGY: linear
    MOLZCULE TYPE: protein
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RESULT 7
US-09-258-643-4
; Sequence 4, Application US/09258643
; Patent No. 6277373
  GENERAL INFORMATION:
    APPLICANT: Hillman, Jennifer L.
    APPLICANT: Lal, Preeti
    APPLICANT: Corley, Neil C.
    APPLICAUT: Shah, Purvi
    TITLE OF INVENTION: NEW HUMAN PHOSPHATIDYLINOSITOL
    NUMBER OF SEQUENCES: 5
    CORRESPONDENCE ADDRESS:
      ADDRESSEE: Incyte Pharmaceuticals, Inc.
      STREET: 3174 Porter Drive
      CITY: Palo Alto
      STATE: CA
      COUNTRY: USA
      ZIP: 94304
    COMPUTER READABLE FORM:
      MEDIUM TYPE: Diskette
      COMPUTER: IBM Compatible
      OPERATING SYSTEM: DOS
      SOFTWARE: FastSEQ for Windows Version 2.0
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/09/258.643
      FILING DATE:
    PRIOR APPLICATION DATA:
      APPLICATION NUMBER: US/08/884,681
      FILING DATE:
    ATTORNEY/AGENT INFORMATION:
      NAME: Billings, Lucy J.
      REGISTRATION NUMBER: 36,749
      REFERENCE/DOCKET NUMBER: PF-0334 US
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: 415-855-0555
      TELEFAX: 415-845-4166
  INFORMATION FOR SEQ ID NO: 4:
    SEQUENCE CHARACTERISTICS:
      LENGTH: 942 amino acids
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TYPE: amino acid
      STRANDEDNESS: single
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                      37.0%; Pred. No. 4.6e-37;
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 Matches 90; Conservative 45; Mismatches 87; Indels
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US-09-418-540-6
; Sequence 6, Application US/09418540
; Patent No. 6296848
  GENERAL INFORMATION:
    APPLICANT: Pot, David A.
    APPLICANT: Williams, Lewis T.
    APPLICANT: Jefferson, Anne Bennett
    APPLICANT: Majerus, Philip W.
    TITLE OF INVENTION: No. 6296848el Grb2 Associating Protein and Nucleic
    TITLE OF INVENTION: Acids Encoding Therefor
    NUMBER OF SEQUENCES: 10
    CORRESPONDENCE ADDRESS:
      ADDRESSEE: Townsend and Townsend and Crew
      STREET: One Market Plaza, Steuart Tower, Suite 2000
      CITY: San Francisco
      STATE: California
      COUNTRY: USA
      ZIP: 94105
    COMPUTER READABLE FORM:
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      COMPUTER: IBM PC compatible
      OPERATING SYSTEM: PC-DOS/MS-DOS
      SOFTWARE: PatentIn Release #1.0, Version #1.25
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      CLASSIFICATION:
    PRIOR APPLICATION DATA:
      APPLICATION NUMBER: US 08/560,005
      FILING DATE: 17-NOV-1995
    ATTORNEY/AGENT INFORMATION:
      NAME: Dow, Karen B.
      REGISTRATION NUMBER: 29,684
      REFERENCE/DOCKET NUMBER: 2307K-0624000
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: 415-326-2400
      TELEFAX: 415-326-2422
  INFORMATION FOR SEQ ID NO: 6:
    SEQUENCE CHARACTERISTICS:
      LENGTH: 942 amino acids
      TYPE: amino acid
      TOPOLOGY: linear
    MOLECULE TYPE: protein
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      NAME/KEY: Region
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     OTHER INFORMATION: /note= "majptase"
US-J9-418-540-6
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US-09-969-528-6
; Sequence 6, Application US/09969528
; Patent No. 6472197
; GENERAL INFORMATION:
       APPLICANT: Pot, David A.
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Williams, Lewis T.
                  Jefferson, Anne Bennett
                  Majerus, Philip W.
        TITLE OF INVENTION: No. 6472197el Grb2 Associating Protein and Nucleic
                          Acids Encoding Therefor
        NUMBER OF SEQUENCES: 10
        CORRESPONDENCE ADDRESS:
             ADDRESSEE: Townsend and Townsend and Crew
             STREET: One Market Plaza, Steuart Tower, Suite 2000
            CITY: San Francisco
             STATE: California
             COUNTRY: USA
             ZIP: 94105
        COMPUTER READABLE FORM:
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             OPERATING SYSTEM: PC-DOS/MS-DOS
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             FILING DATE: 01-Oct-2001
             CLASSIFICATION: <Unknown>
        PRIOR APPLICATION DATA:
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             FILING DATE: <Unknown>
        ATTORNEY/AGENT INFORMATION:
            NAME: Dow, Karen B.
             REGISTRATION NUMBER: 29,684
             REFERENCE/DOCKET NUMBER: 2307K-0624000
        TELECOMMUNICATION INFORMATION:
             TELEPHONE: 415-326-2400
             TELEFAX: 415-326-2422
   INFORMATION FOR SEQ ID NO: 6:
        SEQUENCE CHARACTERISTICS:
             LENGTH: 942 amino acids
             TYPE: amino acid
             TOPOLOGY: linear
        MOLECULE TYPE: protein
        FEATURE:
             NAME/KEY: Region
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             OTHER INFORMATION: /note= "majptase"
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US-09-969-528-6
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RESULT 10
US-08-560-005-7
; Sequence 7, Application US/08560005
; Patent No. 6001354
  GENERAL INFORMATION:
    APPLICANT: Pot, David A.
    APPLICANT: Williams, Lewis T.
    APPLICANT: Jefferson, Anne Bennett
    APPLICANT: Majerus, Philip W.
    TITLE OF INVENTION: No. 6001354el Grb2 Associating Protein and Nucleic
    TITLE OF INVENTION: Acids Encoding Therefor
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    NUMBER OF SEQUENCES: 10
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    CORRESPONDENCE ADDRESS:
      ADDRESSEE: Townsend and Townsend and Crew
      STREET: One Market Plaza, Steuart Tower, Suite 2000
      CITY: San Francisco
      STATE: California
      COUNTRY: USA
      ZIP: 94105
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      FILING DATE:
      CLASSIFICATION: 435
    ATTORNEY/AGENT INFORMATION:
      NAME: Dow, Karen B.
      REGISTRATION NUMBER: 29,684
      REFERENCE/DOCKET NUMBER: 2307K-0624000
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: 415-326-2400
      TELEFAX: 415-326-2422
  INFORMATION FOR SEQ ID NO: 7:
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      TOPOLOGY: linear
    MOLECULE TYPE: protein
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; Sequence 7, Application US/09418540
; Patent No. 6296848
  GENERAL INFORMATION:
    APPLICANT: Pot, David A.
    APPLICANT: Williams, Lewis T.
    APPLICANT: Jefferson, Anne Bennett
    APPLICANT: Majerus, Philip W.
    TITLE OF INVENTION: No. 6296848el Grb2 Associating Protein and Nucleic
    TITLE OF INVENTION: Acids Encoding Therefor
    NUMBER OF SEQUENCES: 10
    CORRESPONDENCE ADDRESS:
     ADDRESSEE: Townsend and Townsend and Crew
     STREET: One Market Plaza, Steuart Tower, Suite 2000
     CITY: San Francisco
     STATE: California
     COUNTRY: USA
      ZIP: 94105
    COMPUTER READABLE FORM:
     MEDIUM TYPE: Floppy disk
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      OPERATING SYSTEM: PC-DOS/MS-DOS
      SOFTWARE: PatentIn Release #1.0, Version #1.25
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APPLICATION NUMBER: US/09/418,540
     FILING DATE: 14-OCT-1999
     CLASSIFICATION:
    PRIOR APPLICATION DATA:
     APPLICATION NUMBER: US 08/560,005
     FILING DATE: 17-NOV-1995
    ATTORNEY/AGENT INFORMATION:
     NAME: Dow, Karen B.
     REGISTRATION NUMBER: 29,684
     REFERENCE/DOCKET NUMBER: 2307K-0624000
    TELECOMMUNICATION INFORMATION:
     TELEPHONE: 415-326-2400
     TELEFAX: 415-326-2422
  INFORMATION FOR SEQ ID NO: 7:
    SEQUENCE CHARACTERISTICS:
     LENGTH: 968 amino acids
     TYPE: amine acid
     TOPOLOGY: linear
    MOLECULE TYPE: protein
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    NAME/KEY: Region
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     OTHER INFORMATION: /note= "ocr1"
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; Sequence 7, Application US/09969528

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; Patent No. 6472197
   GENERAL INFORMATION:
        APPLICANT: Pot, David A.
                   Williams, Lewis T.
                   Jefferson, Anne Bennett
                   Majerus, Philip W.
        TITLE OF INVENTION: No. 6472197el Grb2 Associating Protein and Nucleic
                           Acids Encoding Therefor
        NUMBER OF SEQUENCES: 10
        CORRESPONDENCE ADDRESS:
             ADDRESSEE: Townsend and Townsend and Crew
             STREET: One Market Plaza, Steuart Tower, Suite 2000
             CITY: San Francisco
             STATE: California
             COUNTRY: USA
             ZIP: 94105
        COMPUTER READABLE FORM:
             MEDIUM TYPE: Floppy disk
             COMPUTER: IBM PC compatible
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             CLASSIFICATION: <Unknown>
        PRIOR APPLICATION DATA:
             APPLICATION NUMBER: 08/560,005
             FILING DATE: <Unknown>
        ATTORNEY/AGENT INFORMATION:
             NAME: Dow, Karen B.
             REGISTRATION NUMBER: 29,684
             REFERENCE/DOCKET NUMBER: 2307K-0624000
        TELECOMMUNICATION INFORMATION:
             TELEPHONE: 415-326-2400
             TELEFAX: 415-326-2422
   INFORMATION FOR SEQ ID NO: 7:
        SEQUENCE CHARACTERISTICS:
             LENGTH: 968 amino acids
;
             TYPE: amino acid
             TOPOLOGY: linear
        MOLECULE TYPE: protein
        FEATURE:
             NAME/KEY: Region
             LOCATION: 1..968
             OTHER INFORMATION: /note= "ocr1"
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US-09-969-528-7
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  Best Local Similarity 29.1%; Pred. No. 3.8e-30;
 Matches 102; Conservative 55; Mismatches 121; Indels 73; Gaps
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; Sequence 5, Application US/08884681
; Patent No. 5955338
  GENERAL INFORMATION:
    APPLICANT: Hillman, Jennifer L.
    APPLICANT: Lal, Preeti
    APPLICANT: Corley, Neil C.
    APPLICANT: Shah, Purvi
    TITLE OF INVENTION: NEW HUMAN PHOSPHATIDYLINOSITOL
    NUMBER OF SEQUENCES: 5
    CORRESPONDENCE ADDRESS:
      ADDRESSEE: Incyte Pharmaceuticals, Inc.
     STREET: 3174 Porter Drive
     CITY: Palo Alto
     STATE: CA
     COUNTRY: USA
      ZIP: 94304
    COMPUTER READABLE FORM:
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      FILING DATE: Filed Herewith
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    ATTORNEY/AGENT INFORMATION:
      NAME: Billings, Lucy J.
      REGISTRATION NUMBER: 36,749
      REFERENCE/DOCKET NUMBER: PF-0334 US
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: 415-855-0555
      TELEFAX: 415-845-4166
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; Patent No. 6277373
  GENERAL INFORMATION:
    APPLICANT: Hillman, Jennifer L.
    APPLICANT: Lal, Preeti
   APPLICANT: Corley, Neil C.
    APPLICANT: Shah, Purvi
    WITTE OF INVENTION: NEW HUMAN PHOSPHATIDYLINOSITOL
   NUMBER OF SEQUENCES: 5
    CORRESPONDENCE ADDRESS:
     ADDRESSEE: Incyte Pharmaceuticals, Inc.
     STREET: 3174 Porter Drive
     CITY: Palo Alto
     STATE: CA
     COUNTRY: USA
     ZIP: 94304
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    PRIOR APPLICATION DATA:
     APPLICATION NUMBER: US/08/884,681
     FILING DATE:
    ATTORNEY/AGENT INFORMATION:
     NAME: Billings, Lucy J.
     REGISTRATION NUMBER: 36,749
     REFERENCE/DOCKET NUMBER: PF-0334 US
    TELECOMMUNICATION INFORMATION:
     TELEPHONE: 415-855-0555
     TELEFAX: 415-845-4166
 INFORMATION FOR SEQ ID NO: 5:
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     LENGTH: 901 amino acids
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     LIBRARY: GenBank
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  APPLICANT: VENTER, J. Craig et al.
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  TITLE OF INVENTION: WITH HUMAN DISEASE, METHODS OF DETECTION AND USES
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GenCore version 5.1.6 Copyright (c) 1993 - 2005 Compugen Ltd.

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Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

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3	2012	99.6	478	13	US-10-161-510-11	Sequence 11, Appl
4	2010	99.5	372	13	US-10-161-510-13	Sequence 13, Appl
5	2010	99.5	448	13	US-10-161-510-12	Sequence 12, Appl
6	1791	88.7	329	9	US-09-892-287-3	Sequence 3, Appli
7	872.5	43.2	1001	15	US-10-415-147-3	Sequence 3, Appli
8	868.5	43.0	639	15	US-10-104-047-3268	Sequence 3268, Ap
9	868.5	43.0	639	15	US-10-108-260A-3425	Sequence 3425, Ap
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ALI.GNMENTS

RESULT 1 US-09-892-287-1

[;] Sequence 1, Application US/09892287

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; Patent No. US20020127217A1
   GENERAL INFORMATION:
        APPLICANT: Hillman, Jennifer L.
                  Lal, Preeti
                  Corley, Neil C.
                  Shah, Purvi
        TITLE OF INVENTION: NEW HUMAN PHOSPHATIDYLINOSITOL
                           (4,5) BISPHOSPHATE 5-PHOSPHATASE
        NUMBER OF SEQUENCES: 5
        CORRESPONDENCE ADDRESS:
             ADDRESSEE: Incyte Pharmaceuticals, Inc.
             STREET: 3174 Porter Drive
             CITY: Palo Alto
             STATE: CA
             COUNTRY: USA
             ZIP: 94304
        COMPUTER READABLE FORM:
             MEDIUM TYPE: Diskette
             COMPUTER: IBM Compatible
             OPERATING SYSTEM: DOS
             SOFTWARE: FastSEQ for Windows Version 2.0
        CURRENT APPLICATION DATA:
             APPLICATION NUMBER: US/09/892,287
             FILING DATE: 26-Jun-2001
       PRIOR APPLICATION DATA:
             APPLICATION NUMBER: 09/258,643
             FILING DATE: <Unknown>
        ATTORNEY/AGENT INFORMATION:
             NAME: Billings, Lucy J.
             REGISTRATION NUMBER: 36,749
             REFERENCE/DOCKET NUMBER: PF-0334 US
        TELECOMMUNICATION INFORMATION:
             TELEPHONE: 415-855-0555
             TELEFAX: 415-845-4166
   INFORMATION FOR SEQ ID NO: 1:
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; Publication No. US20020192695A1
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  APPLICANT: EXELIXIS, INC.
  TITLE OF INVENTION: PIBS AS MODIFIERS OF THE p53 PATHWAY AND METHODS OF USE
  FILE REFERENCE: EX02-074C
  CURRENT APPLICATION NUMBER: US/10/161,510
  CURRENT FILING DATE: 2002-06-03
  PRIOR APPLICATION NUMBER: US 60/296,076
  PRIOR FILING DATE: 2001-06-05
  PRIOR APPLICATION NUMBER: US 60/328,605
  PRIOR FILING DATE: 2001-10-10
  PRIOR APPLICATION NUMBER: US 60/357,253
  PRIOR FILING DATE: 2002-02-15
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; Sequence 11, Application US/10161510
; Publication No. US20020192695A1
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  TITLE OF INVENTION: PIBs AS MODIFIERS OF THE p53 PATHWAY AND METHODS OF USE
  FILE REFERENCE: EX02-074C
  CURRENT APPLICATION NUMBER: US/10/161,510
  CURRENT FILING DATE: 2002-06-03
  PRIOR APPLICATION NUMBER: US 60/296,076
  PRIOR FILING DATE: 2001-06-05
  PRIOR APPLICATION NUMBER: US 60/328,605
  PRIOR FILING DATE: 2001-10-10
  PRIOR APPLICATION NUMBER: US 60/357,253
  PRIOR FILING DATE: 2002-02-15
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; Publication No. US20020192695A1
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  FILE REFERENCE: EX02-074C
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  PRIOR FILING DATE: 2001-06-05
  PRIOR APPLICATION NUMBER: US 60/328,605
  PRIOR FILING DATE: 2001-10-10
  PRIOR APPLICATION NUMBER: US 60/357,253
  PRIOR FILING DATE: 2002-02-15
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        121 IEDFGLHFVRESIKNRCYGGLWEKDOLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDY 180
Qу
           121 IEDFGLHFVRESIKNRCYGGLWEKDOLSIAKKHDPLLREFOEGRLLFPPTYKFDRNSNDY 180
Db
        181 DTSEKKRKPAWTDRILWRLKRQPCAGPDTPIPPASHFSLSLRGYSSHMTYGISDHKPVSG 240
Qy
           181 DTSEKKRKPAWTDRILWRLKROPCAGPDTPIPPASHFSLSLRGYSSHMTYGISDHKPVSG 240
Db
QУ
        241 TFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVNDY 300
           241 TFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVNDY 300
Dh
Qу
        301 VSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFQIPPGSL 360
           301 VSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYRNSLRSVVGIRRPFQIPPGSL 360
D'n
        361 REDPLGEAQPQI 372
Qу
           361 REDPLGEAQPQI 372
Db
RESULT 5
US-10-161-510-12
; Sequence 12, Application US/10161510
: Publication No. US20020192695A1
; GENERAL INFORMATION:
  APPLICANT: EXELIXIS, INC.
  TITLE OF INVENTION: PIBS AS MODIFIERS OF THE p53 PATHWAY AND METHODS OF USE
  FILE REFERENCE: EX02-074C
  CURRENT APPLICATION NUMBER: US/10/161,510
  CURRENT FILING DATE: 2002-06-03
  PRIOR APPLICATION NUMBER: US 60/296,076
  PRIOR FILING DATE: 2001-06-05
  PRIOR APPLICATION NUMBER: US 60/328,605
  PRIOR FILING DATE: 2001-10-10
  PRIOR APPLICATION NUMBER: US 60/357,253
  PRIOR FILING DATE: 2002-02-15
  NUMBER OF SEO ID NOS: 14
  SOFTWARE: PatentIn version 3.1
 SEQ ID NO 12
   LENGTH: 448
   TYPE: PRT
   ORGANISM: Homo sapiens
US-10-161-510-12
 Query Match
                            Score 2010; DB 13; Length 448;
                     99.5%;
                            Pred. No. 2.7e-187;
 Best Local Similarity 99.5%;
 Matches 370; Conservative
                          0; Mismatches
                                          2;
                                             Indels
                                                                0;
                                                      0:
                                                         Gaps
          1 MDVLSPLSFIKVSHVRMQGILLLVFAKYQHLPYIQILSTKSTPTGLFGYWGNKGGVNICL 60
Qу
           77 MDVLSPLSFIKVSHVRMQGILLLVFAKYQHLPYIQILSTKSTPTGLFGYWGNKGGVNICL 136
Db
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61 KLYGYYVSIINCHLPPHISNNYORLEHFDRILEMONCEGRDIPNILDHDLIIWFGDMNFR 120
Qу
           137 KLYGYYVSIINCHLPPHISNNYQRLEHFDRILEMQNCEGRDIPNILDHDLIIWFGDMNFR 196
Db
        121 IEDFGLHFVRESIKNRCYGGLWEKDQLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDY 180
Qу
           197 IEDFGLHFVRESIKNRCYGGLWEKDQLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDY 256
Db
        181 DTSEKKRKPAWTDRILWRLKRQPCAGPDTPIPPASHFSLSLRGYSSHMTYGISDHKPVSG 240
Qу
           257 DTSEKKRKPAWTDRILWRLKROPCAGPDTPIPPASHFSLSLRGYSSHMTYGISDHKPVSG 316
Db
QУ
        241 TFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVNDY 300
           317 TFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVNDY 376
Db
        301 VSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFQIPPGSL 360
Qÿ
           377 VSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYRNSLRSVVGIRRPFQIPPGSL 436
Db '
        361 REDPLGEAQPQI 372
Qy
           437 REDPLGEAOPQI 448
Db
RESULT 6
US-09-892-287-3
; Sequence 3, Application US/09892287
; Patent No. US20020127217A1
   GENERAL INFORMATION:
       APPLICANT: Hillman, Jennifer L.
                Lal, Preeti
                Corley, Neil C.
                Shah, Purvi
       TITLE OF INVENTION: NEW HUMAN PHOSPHATIDYLINOSITOL
                        (4,5) BISPHOSPHATE 5-PHOSPHATASE
      NUMBER OF SEQUENCES: 5
       CORRESPONDENCE ADDRESS:
           ADDRESSEE: Incyte Pharmaceuticals, Inc.
           STREET: 3174 Porter Drive
           CITY: Palo Alto
           STATE: CA
           COUNTRY: USA
           ZIP: 94304
       COMPUTER READABLE FORM:
           MEDIUM TYPE: Diskette
           COMPUTER: IBM Compatible
           OPERATING SYSTEM: DOS
           SOFTWARE: FastSEO for Windows Version 2.0
       CURRENT APPLICATION DATA:
           APPLICATION NUMBER: US/09/892,287
           FILING DATE: 26-Jun-2001
       PRIOR APPLICATION DATA:
           APPLICATION NUMBER: 09/258,643
           FILING DATE: <Unknown>
       ATTORNEY/AGENT INFORMATION:
```

```
NAME: Billings, Lucy J.
           REGISTRATION NUMBER: 36,749
           REFERENCE/DOCKET NUMBER: PF-0334 US
       TELECOMMUNICATION INFORMATION:
           TELEPHONE: 415-855-0555
           TELEFAX: 415-845-4166
   INFORMATION FOR SEQ ID NO: 3:
       SEQUENCE CHARACTERISTICS:
           LENGTH: 329 amino acids
           TYPE: amino acid
           STRANDEDNESS: single
           TOPOLOGY: linear
       IMMEDIATE SOURCE:
           LIBRARY: GenBank
           CLONE: 1399101
       SEQUENCE DESCRIPTION: SEQ ID NO: 3:
US-09-892-287-3
 Query Match
                      88.7%;
                            Score 1791; DB 9; Length 329;
 Best Local Similarity
                      99.4%;
                           Pred. No. 4.3e-166;
 Matches 326; Conservative
                            0; Mismatches
                                           2; Indels
                                                          Gaps
                                                                 0;
Qу
         45 GLFGYWGNKGGVNICLKLYGYYVSIINCHLPPHISNNYQRLEHFDRILEMQNCEGRDIPN 104
            Db
          2 GLFGYWGNKGGVNICLKLYGYYVSIINCHLPPHISNNYORLEHFDRILEMONCEGRDIPN 61
        105 ILDHDLIIWFGDMNFRIEDFGLHFVRESIKNRCYGGLWEKDQLSIAKKHDPLLREFQEGR 164
QУ
            62 ILDHDLIIWFGDMNFRIEDFGLHFVRESIKNRCYGGLWEKDOLSIAKKHDPLLREFOEGR 121
Db
        165 LLFPPTYKFDRNSNDYDTSEKKRKPAWTDRILWRLKRQPCAGPDTPIPPASHFSLSLRGY 224
Qу
            Db
        122 LLFPPTYKFDRNSNDYDTSEKKRKPAWTDRILWRLKROPCAGPDTPIPPASHFSLSLRGY 181
        225 SSHMTYGISDHKPVSGTFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPW 284
Qу
            182 SSHMTYGISDHKPVSGTFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPW 241
Db
        285 DWIGLYKVGLRDVNDYVSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYSNSLR 344
Qу
            242 DWIGLYKVGLRDVNDYVSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYRNSLR 301
Db
        345 SVVGISRPFOIPPGSLREDPLGEAOPOI 372
QУ
            302 SVVGIRRPFQIPPGSLREDPLGEAQPQI 329
Db
RESULT 7
US-10-415-147-3
; Sequence 3, Application US/10415147
; Publication No. US20040043399A1
; GENERAL INFORMATION:
  APPLICANT: Bayer AG
  TITLE OF INVENTION: REGULATION OF HUMAN INOSITOL POLYPHOSPHATE 5-PHOSPHATASE
  FILE REFERENCE: LIO235Foreign Countries
  CURRENT APPLICATION NUMBER: US/10/415,147
  CURRENT FILING DATE: 2003-04-25
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PRIOR FILING DATE: 2000-10-30
  PRIOR APPLICATION NUMBER: US 60/257,302
  PRIOR FILING DATE: 2000-12-26
  PRIOR APPLICATION NUMBER: US 60/314,660
  PRIOR FILING DATE: 2001-08-27
  NUMBER OF SEQ ID NOS: 12
  SOFTWARE: PatentIn version 3.1
 SEO ID NO 3
   LENGTH: 1001
   TYPE: PRT
   ORGANISM: Rattus norvegicus
US-10-415-147-3
 Query Match
                      43.2%; Score 372.5; DB 15; Length 1001;
 Best Local Similarity 47.5%; Pred. No. 1.1e-75;
 Matches 169; Conservative 56; Mismatches 126; Indels
                                                      5; Gaps
                                                                4:
          1 MDVLSPLSFIKVSHVRMOGILLLVFAKYOHLPYIOILSTKSTPTGLFGYWGNKGGVNICL 60
QУ:
            483 MDALGPFNFVLVSTVRMQGVILLLFAKYYHLPFLRDVQTDCTRTGLGGYWGNKGGVSVRL 542
D'n
         61 KLYGYYVSIINCHLPPHISNNYORLEHFDRILEMONCEGRDIPNILDHDLIIWFGDMNFR 120
QУ
             Db
        543 AAFGHMLCFLNCHLPAHMDKAEQRKDNFQTILSLQQFQGPGAHGILDHDLVFWFGDLNFR 602
        121 IEDFGLHFVRESIKNRCYGGLWEKDOLSIAKKHDPLLREFOEGRLLFPPTYKFDRNSNDY 180
QУ
            603 IESYDLHFVKFAIDSNOLHOLWEKDOLNMAKNTWPILKGFOEGPLNFAPTFKFDVGTNKY 662
CG
        181 DTSEKKRKPAWTDRILWRLKRQPCAGPDTPIPPASH-FSLSLRGYSSHMTYGISDHKPVS 239
Qy
            Db
        663 DTSAKKRKPAWTDRILWKVK-APSGGP-SPSGRESHRLQVTQHSYRSHMEYTVSDHKPVA 720
        240 GTFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVND 299
QУ
                      Db
        721 ARFLLOFAFRDDVPLVRLEVADEWARPEOAVVRYRVETVFARSSWDWIGLYRVGFRHCKD 780
        300 YVSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFQI 355
Qy
            781 YVAYVWAKHEEV--DGNIYQVTFSEESLPKGHGDFILGYYSHHHSILIGVTEPFQI 834
Db
RESULT 8
US-10-104-047-3268
; Sequence 3268, Application US/10104047
; Publication No. US20030236392A1
; GENERAL INFORMATION:
  APPLICANT: HELIX RESEARCH INSTITUTE
  TITLE OF INVENTION: No. US20030236392A1el full length cDNA
  FILE REFERENCE: H1-A0105
  CURRENT APPLICATION NUMBER: US/10/104,047
  CURRENT FILING DATE: 2002-03-25
  PRIOR APPLICATION NUMBER:
  PRIOR FILING DATE:
  NUMBER OF SEQ ID NOS: 4096
  SOFTWARE: PatentIn Ver. 2.1
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PRIOR APPLICATION NUMBER: US 60/243,745

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LENGTH: 639
   TYPE: PRT
   ORGANISM: Homo sapiens
US-10-104-047-3268
                      43.0%; Score 868.5; DB 15; Length 639;
 Query Match
 Best Local Similarity 47.8%; Pred. No. 1.4e-75;
 Matches 170; Conservative 54; Mismatches 127; Indels
                                                                   4:
          1 MDVLSPLSFIKVSHVRMQGILLLVFAKYQHLPYIQILSTKSTPTGLFGYWGNKGGVNICL 60
Qу
            121 MDALGPFNFVLVSSVRMOGVILLLFAKYYHLPFLRDVOTDCTRTGLGGYWGNKGGVSVRL 180
Db
         61 KLYGYYVSIINCHLPPHISNNYORLEHFDRILEMONCEGRDIPNILDHDLIIWFGDMNFR 120
Qу
              :|: : :||||| |: || ::| || :| :|
                                                 _ | | | | | | | : | | | | | : | | | |
        181 AAFGHMLCFLNCHLPAHMDKAEQRKDNFQTILSLQQFQGPGAQGILDHDLVFWFGDLNFR 240
Db
        121 IEDFGLHFVRESIKNRCYGGLWEKDOLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDY 180
Qу
                             || : ||||: :| :
        241 IESYDLHFVKFAIDSDQLHQLWEKDQLNMAKNTWPILKGFQEGPLNFAPTFKFDVGTNKY 300
ďŒ
        191 DTSEKKRKPAWTDRILWRLKROPCAGPDTPIPPASH-FSLSLRGYSSHMTYGISDHKPVS 239
Qy
            301 DTSAKKRKPAWTDRILWKVK-APGGGP-SPSGRKSHRLQVTQHSYRSHMEYTVSDHKPVA 358
Db
        200 GTFDLELKPLVSAPLIVLMPEDLWTVFNDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVND 299.
QУ
                       359 AOFLLOFAFRDDMPLVRLEVADEWVRPEOAVVPYRMETVFARSSWDWIGLYRVGFRHCKD 418
Db
        300 YVSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFQI 355
QУ
            419 YVAYVWAKHEDV--DGNTYQVTFSEESLPKGHGDFILGYYSHNHSILIGITEPFQI 472
D'n
RESULT 9
US-10-108-260A-3425
; Sequence 3425, Application US/10108260A
; Publication No. US20040005560A1
; GENERAL INFORMATION:
  APPLICANT: HELIX RESEARCH INSTITUTE
  TITLE OF INVENTION: No. US20040005560A1el full length cDNA
  FILE REFERENCE: H1-A0106
  CURRENT APPLICATION NUMBER: US/10/108,260A
  CURRENT FILING DATE: 2002-03-27
  NUMBER OF SEQ ID NOS: 5458
  SOFTWARE: PatentIn Ver. 2.1
 SEQ ID NO 3425
   LENGTH: 639
   TYPE: PRT
   GRGANISM: Homo sapiens
US-10-108-260A-3425
                      43.0%; Score 868.5; DB 15; Length 639;
 Query Match
                      47.8%; Pred. No. 1.4e-75;
 Best Local Similarity
 Matches 170; Conservative 54; Mismatches 127; Indels
                                                                   4;
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: SEO ID NO 3268

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1 MDVLSPLSFIKVSHVRMQGILLLVFAKYQHLPYIQILSTKSTPTGLFGYWGNKGGVNICL 60
Qу
            121 MDALGPFNFVLVSSVRMQGVILLLFAKYYHLPFLRDVQTDCTRTGLGGYWGNKGGVSVRL 180
Db
         61 KLYGYYVSIINCHLPPHISNNYQRLEHFDRILEMQNCEGRDIPNILDHDLIIWFGDMNFR 120
Qу
              :|: : :||||| |: || :| || :| :|
                                                  11111: | 111:11
        181 AAFGHMLCFLNCHLPAHMDKAEQRKDNFQTILSLQQFQGPGAQGILDHDLVFWFGDLNFR 240
Db
        121 IEDFGLHFVRESIKNRCYGGLWEKDOLSIAKKHDPLLREFOEGRLLFPPTYKFDRNSNDY 180
Qу
            241 IESYDLHFVKFAIDSDQLHQLWEKDQLNMAKNTWPILKGFQEGPLNFAPTFKFDVGTNKY 300
Db
        181 DTSEKKRKPAWTDRILWRLKROPCAGPDTPIPPASH-FSLSLRGYSSHMTYGISDHKPVS 239
Qу
            301 DTSAKKRKPAWTDRILWKVK-APGGGP-SPSGRKSHRLQVTQHSYRSHMEYTVSDHKPVA 358
Db
        240 GTFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVND 299
Qy
                       359 AOFILOFAFRDDMPLVRLEVADEWVRPEOAVVRYRMETVFARSSWDWIGLYRVGFRHCKD 418
Db
        300 YVSYAWVGDSKVSCSDNLNOVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFQI 355
Qу
                                 - ::| :|:| ||||:: ::||: |||||
                   419 YVAYVWAKHEDV--DGNTYQVTFSEESLPKGHGDFILGYYSHNHSILIGITEPFQI 472
Db
RESULT 10
US-10-114-270-84
; Sequence 84, Application US/10114270
; Publication No. US20040030110A1
: GENERAL INFORMATION:
  APPLICANT: Guo, Xiaojia
  APPLICANT: Kekuda, Ramesh
  APPLICANT: Miller, Charles E.
  APPLICANT: Malyankar, Uriel M.
  APPLICANT: Spytek, Kimberly A.
  APPLICANT:
             Patturajan, Meera
  APPLICANT: Liu, Ziaohong
  APPLICANT: Gusev, Vladimir Y.
  APPLICANT: Li, Li
  APPLICANT: Vernet, Corine
  APPLICANT: Zerhusen, Bryan D.
  APPLICANT: Gorman, Linda
             Shenoy, Suresh G.
  APPLICANT:
  APPLICANT:
             Pena, Carol E.A.
             Smithson, Glennda
  APPLICANT:
            Burgess, Catherine E.
  APPLICANT:
  APPLICANT: Gerlach, Valerie
  APPLICANT: Padigaru, Muralidhara
  APPLICANT:
             Shimkets, Richard A.
  APPLICANT:
            Gangolli, Esha A.
             Taupier Jr., Raymond J.
  APPLICANT:
  APPLICANT:
             Casman, Stacie J.
  APPLICANT:
            Ji, Weizhen
  APPLICANT: Anderson, David W.
  APPLICANT: Liete, Mario W.
```

Rastelli, Luca

APPLICANT: Edinger, Shlomit R.

APPLICANT:

```
APPLICANT: Stone, David J.
  APPLICANT: MacDougall, John R.
  APPLICANT: Rothenberg, Mark E.
  TITLE OF INVENTION: No. US20040030110A1el Proteins and Nucleic Acids Encoding
  FILE REFERENCE: 21402-322C
  CURRENT APPLICATION NUMBER: US/10/114,270
  CURRENT FILING DATE: 2002-11-27
  PRIOR APPLICATION NUMBER: 60/281,086
  PRIOR FILING DATE: 2001-04-03
  PRIOR APPLICATION NUMBER: 60/281,136
  PRIOR FILING DATE: 2001-04-03
  PRIOR APPLICATION NUMBER: 60/281,863
  PRIOR FILING DATE: 2001-04-05
  PRIOR APPLICATION NUMBER: 60/281,906
  PRIOR FILING DATE: 2001-04-05
  PRIOR APPLICATION NUMBER: 60/282,020
  PRIOR FILING DATE: 2001-04-06
  PRIOR APPLICATION NUMBER: 60/282,930
  PRIOR FILING DATE: 2001-04-10
  PRIOR APPLICATION NUMBER: 60/282,934
  PRIOR FILING DATE: 2001-04-10
  PRIOR APPLICATION NUMBER: 60/283,512
  PRIOR FILING DATE: 2001-04-12
  PRIOR APPLICATION NUMBER: 60/283,710
  PRIOR FILING DATE: 2001-04-13
  PRIOR APPLICATION NUMBER: 60/284,234
  PRIOR FILING DATE: 2001-04-17
  Remaining Prior Application data removed - See File Wrapper or PALM.
 NUMBER OF SEQ ID NOS: 470
 SEQ ID NO 84
   LENGTH: 1006
   TYPE: PRT
   ORGANISM: Homo sapiens
US-10-114-270-84
                       43.0%; Score 868.5; DB 15; Length 1006;
 Query Match
 Best Local Similarity 47.8%; Pred. No. 2.7e-75;
 Matches 170; Conservative 54; Mismatches 127; Indels
                                                         5; Gaps . 4;
          1 MDVLSPLSFIKVSHVRMQGILLLVFAKYQHLPYIQILSTKSTPTGLFGYWGNKGGVNICL 60
Qу
            488 MDALGPFNFVLVSSVRMQGVILLLFAKYYHLPFLRDVQTDCTRTGLGGYWGNKGGVSVRL 547
         61 KLYGYYVSIINCHLPPHISNNYQRLEHFDRILEMQNCEGRDIPNILDHDLIIWFGDMNFR 120
Qу
              548 AAFGHMLCFLNCHLPAHMDKAEQRKDNFQTILSLQQFQGPGAQGILDHDLVFWFGDLNFR 607
Dh
        121 IEDFGLHFVRESIKNRCYGGLWEKDQLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDY 180
Qу
            608 IESYDLHFVKFAIDSDQLHQLWEKDQLNMAKNTWPILKGFQEGPLNFAPTFKFDVGTNKY 667
Db
        181 DTSEKKRKPAWTDRILWRLKROPCAGPDTPIPPASH-FSLSLRGYSSHMTYGISDHKPVS 239
Qу
            668 DTSAKKRKPAWTDRILWKVK-APGGGP-SPSGRKSHRLQVTQHSYRSHMEYTVSDHKPVA 725
Db
        240 GTFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVND 299
QУ
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726 AQFLLQFAFRDDMPLVRLEVADEWVRPEQAVVRYRMETVFARSSWDWIGLYRVGFRHCKD 785
Db
        300 YVSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFQI 355
Qy
            786 YVAYVWAKHEDV--DGNTYQVTFSEESLPKGHGDFILGYYSHNHSILIGITEPFQI 839
Db
RESULT 11
US-10-415-147-12
; Sequence 12, Application US/10415147
; Publication No. US20040043399A1
; GENERAL INFORMATION:
  APPLICANT: Bayer AG
  TITLE OF INVENTION: REGULATION OF HUMAN INOSITOL POLYPHOSPHATE 5-PHOSPHATASE
  FILE REFERENCE: LIO235Foreign Countries
  CURRENT APPLICATION NUMBER: US/10/415,147
  CURRENT FILING DATE: 2003-04-25
  PRIOR APPLICATION NUMBER: US 60/243,745
  PRIOR FILING DATE: 2000-10-30
  PRIOR APPLICATION NUMBER: US 60/257,302
  PRIOR FILING DATE: 2000-12-26
  PRIOR APPLICATION NUMBER: US 60/314,660
  PRIOR FILING DATE: 2001-08-27
  NUMBER OF SEQ ID NOS: 12
  SOFTWARE: PatentIn version 3.1
; SEQ ID NO 12
   LENGTH: 1006
   TYPE: PRT
   ORGANISM: Homo sapiens
US-10-415-147-12
 Query Match
                     43.0%; Score 868.5; DB 15; Length 1006;
 Best Local Similarity
                     47.8%; Pred. No. 2.7e-75;
 Matches 170; Conservative 54; Mismatches 127;
                                              Indels
          1 MDVLSPLSFIKVSHVRMQGILLLVFAKYQHLPYIQILSTKSTPTGLFGYWGNKGGVNICL 60
Qу
            Db
        488 MDALGPFNFVLVSSVRMQGVILLLFAKYYHLPFLRDVQTDCTRTGLGGYWGNKGGVSVRL 547
         61 KLYGYYVSIINCHLPPHISNNYQRLEHFDRILEMQNCEGRDIPNILDHDLIIWFGDMNFR 120
QУ
             548 AAFGHMLCFLNCHLPAHMDKAEQRKDNFQTILSLQQFQGPGAQGILDHDLVFWFGDLNFR 607
Db
        121 IEDFGLHFVRESIKNRCYGGLWEKDQLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDY 180
QУ
            || : ||||: :| :
                            608 IESYDLHFVKFAIDSDOLHOLWEKDOLNMAKNTWPILKGFOEGPLNFAPTFKFDVGTNKY 667
D'n
        181 DTSEKKRKPAWTDRILWRLKRQPCAGPDTPIPPASH-FSLSLRGYSSHMTYGISDHKPVS 239
Qу
            668 DTSAKKRKPAWTDRILWKVK-APGGGP-SPSGRKSHRLQVTQHSYRSHMEYTVSDHKPVA 725
Dρ
        240 GTFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVND 299
QУ
                      | |:
        726 AQFLLQFAFRDDMPLVRLEVADEWVRPEQAVVRYRMETVFARSSWDWIGLYRVGFRHCKD 785
Db
        300 YVSYAWVGDSKVSCSDNLNOVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFOI 355
QУ
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Db

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RESULT 12
US-10-415-147-2
; Sequence 2, Application US/10415147
; Publication No. US20040043399A1
; GENERAL INFORMATION:
  APPLICANT: Bayer AG
  TITLE OF INVENTION: REGULATION OF HUMAN INOSITOL POLYPHOSPHATE 5-PHOSPHATASE
  FILE REFERENCE: LIO235Foreign Countries
  CURRENT APPLICATION NUMBER: US/10/415,147
  CURRENT FILING DATE: 2003-04-25
  PRIOR APPLICATION NUMBER: US 60/243,745
  PRIOR FILING DATE: 2000-10-30
  PRIOR APPLICATION NUMBER: US 60/257,302
  PRIOR FILING DATE: 2000-12-26
  PRIOR APPLICATION NUMBER: US 60/314,660
  PRIOR FILING DATE: 2001-08-27
  NUMBER OF SEQ ID NOS: 12
  SOFTWARE: PatentIn version 3.1
 SEQ ID NO 2
   LENGTH: 751
   TYPE: PRT
   ORGANISM: Homo sapiens
US-10-415-147-2
                       41.2%; Score 831.5; DB 15; Length 751;
 Query Match
 Best Local Similarity 42.9%; Pred. No. 7.2e-72;
 Matches 169; Conservative 56; Mismatches 126; Indels
                                                         43; Gaps
                                                                     5;
          1 MDVLSPLSFIKVSH------
                                                  ----VRMOGILLLVFA 26
Qу
                                                        1 | | | | | : : | | : | |
            || | | :|: |:|
         195 MDALGPFNFVLVTHPSPPGQPETLLNSWLQLYPGSLWGPLGLCGWVSSVRMQGVILLLFA 254
Db
          27 KYQHLPYIQILSTKSTPTGLFGYWGNKGGVNICLKLYGYYVSIINCHLPPHISNNYQRLE 86
Qу
            255 KYYHLPFLRDVQTDCTRTGLGGYWGNKGGVSVRLAAFGHMLCFLNCHLPAHMDKAEQRKD 314
D.b
          87 HFDRILEMQNCEGRDIPNILDHD----LIIWFGDMNFRIEDFGLHFVRESIKNRCYGGLW 142
QУ
                                    | | | | | :
         315 NFQTILSLQQFQGPGAQGILDHEYGLGLVFWFGDLNFRIESYDLHFVKFAIDSDQLHQLW 374
Db
         143 EKDOLSIAKKHDPLLREFOEGRLLFPPTYKFDRNSNDYDTSEKKRKPAWTDRILWRLKRQ 202
Qу
            375 EKDQLNMAKNTWPILKGFQEGPLNFAPTFKFDVGTNKYDTSAKKRKPAWTDRILWKVK-A 433
Db
         203 PCAGPDTPIPPASH-FSLSLRGYSSHMTYGISDHKPVSGTFDLELKPLVSAPLIVLMPED 261
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                      434 PGGGP-SPSGRKSHRLQVTQHSYRSHMEYTVSDHKPVAAQFLLQFAFRDDMPLVRLEVAD 492
Db
         262 LWTVENDMMVSYSSTSDFPSSPWDWIGI,YKVGLRDVNDYVSYAWVGDSKVSCSDNLNQVY 321
Qу
                   493 EWVRPEQAVVRYRMETVFARSSWDWIGLYRVGFRHCKDYVAYVWAKHEDV--DGNTYQVT 550
Db
         322 IDISNIPTTEDEFLLCYYSNSLRSVVGISRPFQI 355
QУ
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RESULT 13
US-10-161-510-10
; Sequence 10, Application US/10161510
; Publication No. US20020192695A1
; GENERAL INFORMATION:
  APPLICANT: EXELIXIS, INC.
  TITLE OF INVENTION: PIBs AS MODIFIERS OF THE p53 PATHWAY AND METHODS OF USE
  FILE REFERENCE: EX02-074C
  CURRENT APPLICATION NUMBER: US/10/161,510
  CURRENT FILING DATE: 2002-06-03
  PRIOR APPLICATION NUMBER: US 60/296,076
  PRIOR FILING DATE: 2001-06-05
  PRIOR APPLICATION NUMBER: US 60/328,605
  PRIOR FILING DATE: 2001-10-10
  PRIOR APPLICATION NUMBER: US 60/357,253
  PRIOR FILING DATE: 2002-02-15
  NUMBER OF SEQ ID NOS: 14
  SOFTWARE: PatentIn version 3.1
; SEQ ID NO 10
   LENGTH: 1056
   TYPE: PRT
   ORGANISM: Homo sapiens
US-10-161-510-10
  Query Match
                       41.2%; Score 831.5; DB 13; Length 1056;
  Best Local Similarity 42.9%; Pred. No. 1.2e-71;
  Matches 169; Conservative 56; Mismatches 126; Indels
                                                         43; Gaps
           1 MDVLSPLSFIKVSH------------------------VRMQGILLLVFA 26
Qу
                                                        | | | | | | : : | | : | |
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         500 MDALGPFNFVLVTHPSPPGOPETLLNSWLQLYPGSLWGPLGLCGWVSSVRMQGVILLLFA 559
Db
          27 KYQHLPYIQILSTKSTPTGLFGYWGNKGGVNICLKLYGYYVSIINCHLPPHISNNYQRLE 86
QУ
             560 KYYHLPFLRDVQTDCTRTGLGGYWGNKGGVSVRLAAFGHMLCFLNCHLPAHMDKAEQRKD 619
Db
          87 HFDRILEMQNCEGRDIPNILDHD----LIIWFGDMNFRIEDFGLHFVRESIKNRCYGGLW 142
Ov
             :| || :| :|
                             620 NFQTILSLQQFQGPGAQGILDHEYGLGLVFWFGDLNFRIESYDLHFVKFAIDSDQLHQLW 679
Db
         143 EKDOLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDYDTSEKKRKPAWTDRILWRLKRQ 202
QУ
             680 EKDQLNMAKNTWPILKGFQEGPLNFAPTFKFDVGTNKYDTSAKKRKPAWTDRILWKVK-A 738
DЪ
         203 PCAGPDTPIPPASH-FSLSLRGYSSHMTYGISDHKPVSGTFDLELKPLVSAPLIVLMPED 261
Qу
                     |\cdot|: |
         739 PGGGP-SPSGRKSHRLQVTQHSYRSHMEYTVSDHKPVAAQFLLQFAFRDDMPLVRLEVAD 797
Dω
         262 LWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGI,RDVNDYVSYAWVGDSKVSCSDNLNQVY 321
QУ
                    Db
         798 EWVRPEQAVVRYRMETVFARSSWDWIGLYRVGFRHCKDYVAYVWAKHEDV--DGNTYQVT 855
         322 IDISNIPTTEDEFLLCYYSNSLRSVVGISRPFQI 355
Qу
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RESULT 14
US-10-106-698-4755
; Sequence 4755, Application US/10106698
: Publication No. US20030109690A1
; GENERAL INFORMATION:
  APPLICANT: Ruben et al.
  TITLE OF INVENTION: Colon and Colon Cancer Associated Polynucleotides and
Polypeptides
  FILE REFERENCE: PA005P1
  CURRENT APPLICATION NUMBER: US/10/106,698
  CURRENT FILING DATE: 2002-03-27
  PRIOR APPLICATION NUMBER: PCT/US00/26524
  PRIOR FILING DATE: 2000-09-28
  PRIOR APPLICATION NUMBER: US 60/157,137
  PRIOR FILING DATE: 1999-09-29
  PRIOR APPLICATION NUMBER: US 60/163,280
  PRIOR FILING DATE: 1999-11-03
  NUMBER OF SEQ ID NOS: 8564
  SOFTWARE: PatentIn Ver. 3.0
; SEQ ID NO 4755
   LENGTH: 279
   TYPE: PRT
   ORGANISM: Homo sapiens
   FEATURE:
   NAME/KEY: MISC FEATURE
   LOCATION: (247)
   OTHER INFORMATION: Xaa equals any of the naturally occurring L-amino acids
US-10-106-698-4755
                       33.8%; Score 683.5; DB 14; Length 279;
 Query Match
 Dest Local Similarity 52.8%; Pred. No. 5.1e-58;
 Matches 130; Conservative 39; Mismatches 74;
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                                                                     3;
           1 MDVLSPLSFIKVSHVRMQGILLLVFAKYQHLPYIQILSTKSTPTGLFGYWGNKGGVNICL 60
Qу
            10 MDALGPFNFVLVSSVRMOGVILLLFAKYYHLPFLRDVQTDCTRTGLGGYWGNKGGVSVRL 69
Db
          61 KLYGYYVSIINCHLPPHISNNYORLEHFDRILEMONCEGRDIPNILDHDLIIWFGDMNFR 120
Οv
              :|: : :||||| |:
                               70 AAFGHMLCFLNCHLPAHMDKAEQRKDNFQTILSLQQFQGPQAQGILDHDLVFWFGDLNFR 129
Db
         121 IEDFGLHFVRESIKNRCYGGLWEKDOLSIAKKHDPLLREFOEGRLLFPPTYKFDRNSNDY 180
Çу
                               || : ||||: :| :
         130 IESYDLHFVKFAIDSDQLHQLWEKDQLNMAKNTWPILKGFQEGPLNFAPTFKFDVGTNKY 189
Db
         181 DTSEKKRKPAWTDRILWRLKRQPCAGPDTPIPPASH-FSLSLRGYSSHMTYGISDHKPVS 239
ÒЯ
            :: | | | | | | : | | | | |
Db
         190 DTSAKKRKPAWTDRILWKVK-APGGGP-SPSGRKSHRLQVTQHSYRSHMEYTVSDHKPVX 247
         240 GTFDLE 245
Qу
              1 1:
Db
         248 AQFLLO 253
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RESULT 15
US-10-264-049-4157
; Sequence 4157, Application US/10264049
; Publication No. US20040005579A1
; GENERAL INFORMATION:
; APPLICANT: Birse et al.
  TITLE OF INVENTION: Nucleic Acids, Proteins, and Antibodies
  FILE REFERENCE: PA133P1
  CURRENT APPLICATION NUMBER: US/10/264,049
  CURRENT FILING DATE: 2002-10-04
  PRIOR APPLICATION NUMBER: PCT/US01/18569
  PRIOR FILING DATE: 2001-06-07
 PRIOR APPLICATION NUMBER: US 60/209,467
  PRIOR FILING DATE: 2000-06-07
  NUMBER OF SEQ ID NOS: 4360
  SOFTWARE: PatentIn Ver. 3.1
; SEQ ID NO 4157
   LENGTH: 162
   TYPE: PRT
   ORGANISM: Homo sapiens
   FEATURE:
   NAME/KEY: MISC FEATURE
   LOCATION: (120)
   OTHER INFORMATION: Xaa equals any of the twenty naturally occurring L-amino
acids
   FEATURE:
   NAME/KEY: MISC FEATURE
   LOCATION: (130)
   OTHER INFORMATION: Xaa equals any of the twenty naturally occurring L-amino
US-10-264-049-4157
                      26.8%; Score 541.5; DB 15; Length 162;
 Query Match
 Best Local Similarity 74.6%; Pred. No. 1.7e-44;
 Matches 106; Conservative 0; Mismatches 7; Indels 29; Gaps
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          1 MDVLSPLSFIKVSHVRMQGILLLVFAKYQHLPYIQILSTKSTPTGLFGYWGNKGGVNICL 60
Qv
            26 MDVLSPLSFIKVSHVRMQGILLLVFAKYQHLPYIQILSTKSTPTGLFGYWGNKGGVNICL 85
Db
         61 KLYGYYVSIINCHLPPHISN------NYORLEHFDRILEMONCEGRDIPNII, 106
Qу
            86 KLYGYYVSIINCHLPPHISTITSGWSTLTGSWRCXIVRARH------PNIX 130
Db
        107 DHDLIIWFGDMNFRIEDFGLHF 128
Qу
            Db
         131 DHDLIIWFGDMNFRIEDFGLHF 152
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Search completed: February 10, 2005, 18:05:21 Job time: 135 secs

> GenCore version 5.1.6 Copyright (c) 1993 - 2005 Compugen Ltd.

OM protein - protein search, using sw model

Run on: February 10, 2005, 17:06:26; Search time 41 Seconds

(without alignments)

872.991 Million cell updates/sec

Title: US-09-892-287-1

Perfect score: 2020

Sequence: 1 MDVLSPLSFIKVSHVRMQGI......FQIPPGSLREDPLGEAQPQI 372

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 283416 seqs, 96216763 residues

Total number of hits satisfying chosen parameters: 283416

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : PIR_79:*

1: pir1:*

2: pir2:*

3: pir3:*

4: pir4:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

		ક				
Result `		Query	_			
No.	Score	Match	Length	DB	ID	Description
1	406.5	20.1	1017	2	T42384	inositol-1,4,5-tri
2	367.5	18.2	893	2	G59431	phosphatidylinosit
3	367.5	18.2	970	2	S29069	inositol polyphosp
4	334	16.5	672	2	A41075	inositol-1,4,5-tri
5	334	16.5	1216	2	JW0105	synaptojanin 2 alp
6	328	16.2	1575	2	368448	synaptojanin, 170K
7	300.5	15.3	1136	2	D86190	hypothetical prote
3	306	15.1	1305	2	T00670	probable inositol
9	300.5	14.9	753	2	T19338	hypothetical prote
10	298.5	14.8	331	2	D96515	hypothetical prote
11	297.5	14.7	401	2	G84792	hypothetical prote
12	291.5	14.4	1107	2	S61667	probable membrane
13	289.5	14.3	946	2	S48433	inositol-1,4,5-tri
14	284	14.1	1144	2	F84725	probable inositol
15	284	14.1	1258	2	JC5765	inositol polyphosp
16	279.5	13.8	1076	2	T40141	phosphatidylinosit
17	279	13.8	501	2	H84727	probable inositol
18	278.5	13.8	590	2	C86465	probable inositol

19	278	13.8	1183	2	T39233	probable Inositol
20	277.5	13.7	574	2	T48113	inositol-1,4,5-tri
21	276	13.7	889	2	T40557	probable inositol
22	268.5	13.3	427	2	C88883	protein JC8.10 [im
23	265.5	13.1	417	2	E84430	probable inositol
24	265	13.1	1183	2	S63046	probable membrane
25	264	13.1	613	2	T51938	inositol-1,4,5-tri
26	264	13.1	670	2	D96739	hypothetical prote
27	263	13.0	595	2	T05087	hypothetical prote
28	260	12.9	646	2	T51937	probable inositol-
29	251	12.4	1189	2	JC6118	SH2-containing ino
30	250	12.4	1188	2	JC4889	phosphatidylinosit
31	240	11.9	993	2	G96680	F5I14.11 [imported
32	227.5	11.3	398	2	S44627	C50C3.7 protein -
33	218	10.8	342	2	T19021	probable inositol
34	172.5	8.5	384	2	S66758	probable membrane
35	142.5	7.1	396	2	T15465	hypothetical prote
36	114	5.6	743	2	S19437	hypothetical prote
37	104.5	5.2	412	2	\$45721	inositol-1,4,5-tri
38	104.5	5.2	446	2	A56733	nuclear domain 10
39	102.5	5.1	412	2	S44357	inositol 1,4,5-tri
40	99	4.9	1364	2	T40839	hypothetical prote
41	93	4.9	384	2	S76402	hypothetical prote
42	97.5	4.8	363	2	A54167	inositol-polyphosp
43	95 . 5	4.7	954	2	S46105	glucan 1,4-alpha-g
44	94.5	4.7	371	2	AE1294	hypothetical prote
45	93	4.6	381	2	T28378	ORF MSV217 SCG gen

ALIGNMENTS

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RESULT 1
T42384
inositol-1,4,5-trisphosphate 5-phosphatase (EC 3.1.3.56) II, isoform 115K,
membrane-associated - mouse
N; Alternate names: inositol trisphosphate phosphomonoesterase
C; Species: Mus musculus (house mouse)
C;Date: 03-Dec-1999 #sequence revision 03-Dec-1999 #text_change 09-Jul-2004
C; Accession: T42384
R; Matzaris, M.; O'Malley, C.; Badger, A.; Speed, C.; Bird, P.I.; Mitchell, C.A.
submitted to the EMBL Data Library, December 1997
A; Description: Distinct membrane and cytosolic forms of inositol polyphosphate
5-phosohatase II. Efficient membrane localization requires two discrete domains.
A; Reference number: Z22143
A; Accession: T42384
A; Status: preliminary; translated from GB/EMBL/DDBJ
A; Molecule type: mRNA
A; Residues: 1-1017 <MAT>
A; Cross-references: UNIPROT:054996; EMBL: AF040094; NID: g2766528; PID: g2766529;
PIDN:AAB95412.1
A; Experimental source: brain
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A;Gene: INPP5P
C; Keywords: phosphoric monoester hydrolase
                          20.1%; Score 406.5; DB 2; Length 1017;
  Query Match
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Best Local Similarity 34.1%; Pred. No. 1.7e-26;
  Matches
           86; Conservative 42; Mismatches 89; Indels
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                                                                        4;
           9 FIKVSHVRMQGILLLVFAKYQHLPYIQILSTKSTPTGLFG----YWGNKGGVNICLKLYG 64
Qу
             415 YAKVKFVRLVGIMLLLYVKQEHAAYISEVEAETVGTGFMGRMSFQQGNKGGVAIRFQLHN 474
Db
          65 YYVSIINCHLPPHISNNYORLEHFDRILEMONCEGRDIP-----NILDHDLIIWFG 115
Qу
                            :|::|
                                                          | ||:|:|
Db
         475 TSICVVNSHLAAHTEEYERRNODYRDI-----CSRMOFPQVDPSQPPLTINKHDVILWLG 529
Qу
         116 DMNFRIEDFGLHFVRESIKNRCYGGLWEKDQLSIAKKHDPLLREFQEGRLLFPPTYKFDR 175
             Db
         530 DLNYRIEELDVGKVKKLVEEKAFQTLYAHDQLKIQVAARTIFDGFTEGEITFQPTYKYDT 589
         176 NSNDYDTSEKKRKPAWTDRILWRLKRQPCAGPDTPIPPASHFSLSLRGYSSHMTYGISDH 235
Qу
              Db
         590 GSDDWDTSESARAPAWCDRILWKGK-------NITQLSYQSHMALKTSDH 632
         236 KPVSGTFDLELK 247
Qу
             Db
         633 KPVSSVFDIGVR 644
RESULT 2
G59431
phosphatidylinositol polyphosphate 5-phosphotase, isoform b [imported] - human
C; Species: Homo sapiens (man)
C;Date: 03-Jun-2002 #sequence revision 03-Jun-2002 #text change 23-Sep-2002
C; Accession: G59431; H59431; A59443; B59432; C59432; D59432; E59432; F59432
R; Duden, R.; Griffiths, G.; Frank, R.; Argos, P.; Kreis, T.E.
Cell 64, 649-665, 1991
A; Title: Beta-COP, a 110 kd protein associated with non-clathrin-coated vesicles
and the Golgi complex, shows homology to beta-adaptin.
A; Reference number: G59431; MUID: 91121521; PMID: 1840503
A; Accession: G59431
A; Status: preliminary
A; Molecule type: DNA
A; Residues: 1-893 < DUD>
A;Cross-references: GB:NP 001578; PID:g13325070; PIDN:NP 001578.2
R; Mueller, O.T.; Hartsfield, J.K. Jr.; Gallardo, L.A.; Essig, Y.P.; Miller,
K.L.; Papenhausen, P.R.; Tedesco, T.A.
Am. J. Hum. Genet. 49, 804-810, 1991
A; Title: Lowe oculocerebrorenal syndrome in a female with a balanced X; 20
translocation: mapping of the X chromosome breakpoint.
A; Reference number: H59431
A; Accession: H59431
A; Status: preliminary
A; Molecule type: DNA
A; Residues: 1-893 < MUE>
A;Cross-references: GB:NP_001578; PID:g13325070; PIDN:NP_001578.2
R; Attree, G.; Olivos, I.M.; Okabe, I.; Bailey, L.C.; Nelson, D.L.; Lewis, R.A.;
McInnes, R.R.; Nussbaum, R.L.
Nature 358, 239-242, 1992
A; Title: The Lowe's oculocerebrorenal syndrome gene encodes a protein highly
homologous to inositol polyphosphate-5-phosphatase.
A; Reference number: A59443; MUID: 92334430; PMID: 1321346
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A; Accession: A59443
A; Status: preliminary
A; Molecule type: DNA
A; Residues: 1-893 < ATT>
A; Cross-references: GB:NP 001578; PID:q13325070; PIDN:NP 001578.2
R; Leahey, A.M.; Charnas, L.R.; Nussbaum, R.L.
Hum. Mol. Genet. 2, 461-463, 1993
A; Title: Nonsense mutations in the OCRL-1 gene in patients with the
oculocerebrorenal syndrome of Lowe.
A; Reference number: B59432; MUID: 93278398; PMID: 8504307
A; Accession: B59432
A; Status: preliminary
A; Molecule type: DNA
A; Residues: 1-893 < LEA>
A; Cross-references: GB:NP 001578; PID:g13325070; PIDN:NP_001578.2
R; Zhang, X.; Jefferson, A.B.; Auethavekiat, V.; Majerus, P.W.
Proc. Natl. Acad. Sci. U.S.A. 92, 4853-4856, 1995
A; Title: The protein deficient in Lowe syndrome is a phosphatidylinositol-4,5-
bisphosphate 5-phosphatase.
A; Reference number: C59432
A; Accession: C59432
A; Status: preliminary
A; Molecule type: DNA
A; Residues: 1-893 < ZHA>
A;Cross-references: GB:NP_001578; PID:g13325070; PIDN:NP_001578.2
R; Nussbaum, R.L.; Orrison, B.M.; Janne, P.A.; Charnas, L.; Chinault, A.C.
Hum. Genet. 99, 145-150, 1997
A; Title: Physical mapping and genomic structure of the Lowe syndrome gene OCRL1.
A; Reference number: D59432
A; Accession: D59432
A; Status: preliminary
A; Molecule type: DNA
A; Residues: 1-893 < NUS>
A; Cross-references: GB:NP 001578; PID:g13325070; PIDN:NP 001578.2
R; Janne, P.A.; Suchy, S.F.; Bernard, D.; MacDonald, M.; Crawley, J.; Grinberg,
A.; Wynshaw-Boris, A.; Westphal, H.; Nussbaum, R.L.
J. Clin. Invest. 101, 2042-2053, 1998
A; Title: Functional overlap between murine Inpp5b and Ocrl1 may explain why
deficiency of the murine ortholog for OCRL1 does not cause Lowe syndrome in
A; Reference number: E59432
A; Accession: E59432
A; Status: preliminary
A, Molecule type: DNA
A; Residues: 1-893 < JAN>
A;Cross-references: GB:NP_001578; PID:g13325070; PIDN:NP_001578.2
R; Erneux, C.; Govaerts, C.; Communi, D.; Pesesse, X.
Biochim. Biophys. Acta 1436 (1-2), 185-199, 1998
A; Title: The diversity and possible functions of the inositol polyphosphate 5-
phosphatases.
A; Reference number: F59432
A; Accession: F59432
A; Status: preliminary
A; Molecule type: DNA
A; Residues: 1-893 < ERN>
A; Cross-references: GB:NP 001578; PID:q13325070; PIDN:NP 001578.2
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Qу
               307 SKAKYKKVQLVRLVGMMLLIFARKDQCRYIRDIATETVGTGIMGKMGNKGGVAVRFVFHN 366
Db
         65 YYVSIINCHLPPHISNNYORLEHFDRILEMON--CEGRDIP--NILDHDLIIWFGDMNFR 120
Qу
                367 TTFCIVNSHLAAHVEDFERRNODYKDICARMSFVVPNOTLPOLNIMKHEVVIWLGDLNYR 426
Db
        121 IEDFGLHFVRESIKNRCYGGLWEKDQLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDY 180
QУ
            427 LCMPDANEVKSLINKKDLORLLKFDOLNIORTOKKAFVDFNEGEIKFIPTYKYDSKTDRW 486
Db
        181 DTSEKKRKPAWTDRILWRLKROPCAGPDTPIPPASHFSLSLRGYSSHMTYGISDHKPVSG 240
Qу
            ::: | | | | |
        437 DSSGKCRVPAWCDRILWR------GTNVNQLNYRSHMELKTSDHKPVSA 529
Db
        241 TFDLELKPL-----VSAPLIVLMPEDLWTVENDMMVSYS-STSDFPSSPWDWIGLYKV 292
Qу
             530 LFHIGVKVVDERRYRKVFEDSVRIMDR----MENDFLPSLELSRREF----- 572
Db
        200 GLRDVNDYVSYAWVGDSKVSCSDNLNQVYIDISNIPTTED 332
Оv
                Db
         573 ----VFENVKFRQLQKEKFQISNN-GQVPCHFSFIPKLND 607
RESULT 3
S29069
inositol polyphosphate-5-phosphatase homolog - human
C; Species: Homo sapiens (man)
C;Date: 25-Jeb-1994 #sequence revision 10-Nov-1995 #text change 09-Jul-2004: 11
C; Accession: S29069; I68621
R; Attree, O.; Olivos, I.M.; Okabe, I.; Bailey, L.C.; Nelson, D.L.; Lewis, R.A.;
McInnes, R.R.; Nussbaum, R.L.
Nature 358, 239-242, 1992
A; Title: The Lowe's oculocerebrorenal syndrome gene encodes a protein highly
homologous to inositol polyphosphate-5-phosphatase.
A; Reference number: S29069; MUID: 92334430; PMID: 1321346
A; Accession: S29069
A; Status: preliminary
A; Molecule type: mRNA
A; Residues: 1-970 <ATT>
A; Cross-references: UNIPROT:Q01968
R; Leahey, A.M.; Charnas, L.R.; Nussbaum, R.L.
Hum. Mol. Genet. 2, 461-463, 1993
A; Title. Noncense mutations in the OCRL-1 gene in patients with the
oculocerebrorenal syndrome of Lowe.
A; Reference number: I54349; MUID: 93278398; PMID: 8504307
A; Accession: I68621
A;Status: preliminary; translated from GB/EMBL/DDBJ
A; Molecule type: mRNA
A; Residues: 383-912 < RES>
A; Cross-references: GB:S62085; NID:q385336; PIDN:AAB26926.1; PID:q385337
C; Genetics:
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A; Gene: OCRL-1
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18.2%; Score 367.5; DB 2; Length 970;
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 Matches 100; Conservative 58; Mismatches 131; Indels
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                                                                    8;
           5 SPLSFIKVSHVRMQGILLLVFAKYQHLPYIQILSTKSTPTGLFGYWGNKGGVNICLKLYG 64
Qу
            384 SKAKYKKVQLVRLVGMMLLIFARKDQCRYIRDIATETVGTGIMGKMGNKGGVAVRFVFHN 443
Db
          65 YYVSIINCHLPPHISNNYQRLEHFDRILEMQN--CEGRDIP--NILDHDLIIWFGDMNFR 120
Qy
                444 TTFCIVNSHLAAHVEDFERRNODYKDICARMSFVVPNQTLPQLNIMKHEVVIWLGDLNYR 503
Db
         121 IEDFGLHFVRESIKNRCYGGLWEKDQLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDY 180
Qy
                 : |: | : | : ||:| :
                                              :| || : | ||||:| :: :
         504 LCMPDANEVKSLINKKDLQRLLKFDQLNIQRTQKKAFVDFNEGEIKFIPTYKYDSKTDRW 563.
Db
         181 DTSEKKRKPAWTDRILWRLKROPCAGPDTPIPPASHFSLSLRGYSSHMTYGISDHKPVSG 240
Qу
            1:1 | 1 | 1 | 1 | 1 | 1 | 1 | 1
                                             :::
                                                   564 DSSGKCRVPAWCDRILWR------GTNVNQLNYRSHMELKTSDHKPVSA 606
Db
         241 TFDLELKPL-----VSAPLIVLMPEDLWTVENDMMVSYS-STSDFPSSPWDWIGLYKV 292
QУ
             507 LFHIGVKVVDERRYRKVFEDSVRIMDR----MENDFLPSLELSRREF------ 649
Db
         293 GLRDVNDYVSYAWVGDSKVSCSDNLNOVYIDISNIPTTED 332
QУ
                1:1::1:
                                        650 ----VFENVKFRQLQKGKFQISNN-GQVPCHFSFIPKLND 684
Db
RESULT 4
A41075
inositol-1,4,5-trisphosphate 5-phosphatase (EC 3.1.3.56) - human (fragment)
C; Species: Homo sapiens (man)
C;Date: 12-Jun-1992 #sequence_revision 12-Jun-1992 #text_change 29-Aug-1997
C; Accession: A41075
R; Ross, T.S.; Jefferson, A.B.; Mitchell, C.A.; Majerus, P.W.
J. Biol. Chem. 266, 20283-20289, 1991
A; Title: Cloning and expression of human 75-kDa inositol polyphosphate-5-
phosphatase.
A; Reference number: A41075; MUID: 92041857; PMID: 1718960
A; Accession: A41075
A; Status: preliminary
A; Molecule type: mRNA
A; Residues: 1-672 < ROS>
A; Cross-references: GB:M74161
C; Genetics:
A;Gene: GDB:INPP5B
A; Cross-references: GDB:129756; OMIM:147264
A; Map position: 1p34-1p34
C; Keywords: phosphoric monoester hydrolase
                       16.5%; Score 334; DB 2; Length 672;
  Query Match
                       31.7%; Pred. No. 1.6e-20;
  Best Local Similarity
  Matches 78; Conservative 45; Mismatches 97; Indels 26; Gaps
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9 FIKVSHVRMQGILLLVFAKYQHLPYIQILSTKSTPTGLFGYWGNKGGVNICLKLYGYYVS 68
Qу
            133 YAKVKLIRLVGIMLLLYVKQEHAAYISEVEAETVGTGIMGRMGNKGGVAIRFQFHNTSIC 192
Db
         69 IINCHLPPHISNNYORLEHF-DRILEMONCE-GRDIP--NILDHDLIIWFGDMNFRIEDF 124
Qу
            193 VVNSHLAAHIEEYERRNQDYKDICSRMQFCQPDPSLPPLTISNHDVILWLGDLNYRIEEL 252
Db
        125 GLHFVRESIKNRCYGGLWEKDOLSIAKKHDPLLREFOEGRLLFPPTYKFDRNSNDYDTSE 184
Qу
            253 DVEKVKKLIEEKDFOMLYAYDOLKIQVAAKTVFEGFTEGELTFQPTYKYDTRA--LTTGI 310
Db
        185 KKRKPAWTDRI---LWRLKROPCAGPDTPIPPASHFSLSLRGYSSHMTYGISDHKPVSGT 241
Qy
                : | | | |
                                         ::: | | | | |
        311 PVRSAVLLPGVIGFLWKGK------NITQLSYQSHMALKTSDHKPVSSV 353
Db
Qу
        242 FDLELK 247
            ||: ::
        354 FDIGVR 359
Db
RESULT 5
JW0105
synaptojanin 2 alpha protein - mouse
C: Species: Mus musculus (house mouse)
C;Date: 10-Jul-1998 #sequence revision 10-Jul-1998 #text change 09-Jul-2004
C; Accession: JW0105
R; Seet, L.F.; Cho, S.; Hessel, A.; Dumont, D.J.
Biochem. Biophys. Res. Commun. 247, 116-122, 1998
A; Title: Molecular cloning of multiple isoforms of synaptojanin 2 and assignment
of the gene to mouse chromosome 17A2-3.1.
A; Reference number: JW0105; MUID: 98300294; PMID: 9636665
A; Contents: Liver
A; Accession: JW0105
A; Molecule type: mRNA
A; Residues: 1-1216 <SEE>
A;Cross-references: UNIPROT:Q9D2G5; GB:AF041862; NID:g3241994; PIDN:AAC40146.1;
PID:g3241995
 Query Match
                      16.5%; Score 334; DB 2; Length 1216;
                      31.2%; Pred. No. 3.7e-20;
 Best Local Similarity
 Matches 92; Conservative 49; Mismatches 124; Indels
                                                      30; Gaps 9;
          9 FIKVSHVRMOGILLLVFAKYQHLPYIQILSTKSTPTGLFGYWGNKGGVNICLKLYGYYVS 68
Qу
            536 YILLTSAOLVGVCLYIFVRPYHVPFIRDVAIDTVKTGMGGKAGNKGAVGIRFQLHSTSFC 595
Db
         69 IINCHLPPHISNNYQRLEHFDRIL-EMQNCEGRDIPNILDHDLIIWFGDMNFRIEDFGLH 127
Qу
                   :| | : | :: | |
                                         Db
        596 FVCSHLTAGOSOVKERNEDYREITHKLSFPSGR---NIFSHDYVFWCGDFNYRI-DLTYE 651
        128 FVRESIKNRCYGGLWEKDQLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDYDTSEKKR 187
QУ
               652 EVFYFVKRQDWKKLMEFDQLQKSSGKIFKDFHEGAVNFGPTYKYDVGSAAYDTSDKCR 711
Db
        188 KPAWTDRILWRLKRQP---CAGP----DTPI---PPASH-FSLSLRGYSSHMTYGISDHK 236
QУ
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712 TPAWTDRVLWWRKKHPYDKTAGELNLLDSDLDGDPQIRHTWSPGTLKYYGRAELQASDHR 771
Db
         237 PVSGTFDLELKPL------VSAPLIVLMPEDLWTVENDMMVSYSSTSDFP 280
Qу
             || ::|::::
                                   772 PVLAIVEVEVQEVDVGARERVFQEVSS---VQGPLDATVVVNLQSPTLEEKNEFP 823
Db
RESULT 6
S68448
synaptojanin, 170K - rat
C; Species: Rattus norvegicus (Norway rat)
C;Date: 17-Jul-1998 #sequence_revision 17-Jul-1998 #text_change 05-Nov-1999
C; Accession: S68448; S78547; S78527
R;McPherson, P.S.; Garcia, E.P.; Slepnev, V.I.; David, C.; Zhang, X.; Grabs, D.;
Sossin, W.S.; Bauerfeind, R.; Nemoto, Y.; de Camilli, P.
Nature 379, 353-357, 1996
A; Title: A presynaptic inositol-5-phosphatase.
A; Reference number: A58183; MUID: 96149250; PMID: 8552192
A; Accession: S68448
A; Status: nucleic acid sequence not shown
A; Molecule type: mRNA
A; Residues: 1-1575 <MCP>
A; Cross-references: EMBL: U45479; NID: g1166574; PIDN: AAB60525.1; PID: g1166575;
PID:q1166576
A; Experimental source: brain
A; Accession: S78547
A; Molecule type: protein
A; Residues: 1075-1093; 1173-1199; 1290-1308 < MCS>
A; Experimental source: brain
R; de Camilli, P.
submitted to the EMBL Data Library, January 1996
A; Reference number: $78527
A: Accession: S78527
A; Molecule type: mRNA
A: Residues: 1-587, 'D', 589-1575 < DEC>
A; Cross-references: EMBL:U45479
C; Function:
A; Description: binds the SH3 domain of amphiphysin
C; Keywords: alternative splicing; alternative termination; synaptic vesicle
F;1-1575/Product: synaptojanin, long form b #status predicted <MAT3>
F;1-1139,1156-1575/Product: synaptojanin, long form a #status predicted <MAT2>
F:1-113,1156-1308/Product: synaptojanin, short form #status predicted <MAT1>
                        16.2%; Score 328; DB 2; Length 1575;
  Query Match
  Best Local Similarity 33.6%; Pred. No. 1.7e-19;
          84; Conservative 44; Mismatches 106; Indels 16; Gaps
 Matches
           9 FIKVSHVRMQGILLLVFAKYQHLPYIQILSTKSTPTGLFGYWGNKGGVNICLKLYGYYVS 68
QУ
             625 YVLLASEQLVGVCLFVFIRPQHAPFIRDVAVDTVKTGMGGATGNKGAVAIRMLFHTTSLC 684
Db
          69 IINCHLPPHISNNYQRLEHFDRIL-EMQNCEGRDIPNILDHDLIIWFGDMNFRIEDFGLH 127
\tilde{\mathbf{C}}\mathbf{\lambda}
                     685 FVCSHFAAGQSQVKERNEDFVEIARKLSFPMGR---MLFSHDYVFWCGDFNYRI-DLPNE 740
Db
         128 FVRESIKNRCYGGLWEKDQLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDYDTSEKKR 187
Oν
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Db
         741 EVKELIRQQNWDSLIAGDQLINQKNAGQIFRGFLEGKVTFAPTYKYDLFSEDYDTSEKCR 800
         188 KPAWTDRILWRLKRQPC--AGPDTPIPPASHFSLSLRGYS----SHMTYG-----ISDHK 236
Qу
               801 TPAWTDRVLWRRRKWPFDRSAEDLDLLNASFODESKILYTWTPGTLLHYGRAELKTSDHR 860
Db
         237 PVSGTFDLEL 246
QУ
             | | | :::
         861 PVVALIDIDI 870
Db
RESULT 7
D86190
hypothetical protein [imported] - Arabidoosis thaliana
C; Species: Arabidopsis thaliana (mouse-ear cress)
C;Date: 02-Mar-2001 #sequence revision 02-Mar-2001 #text change 09-Jul-2004
C; Accession: D86190
R; Theologis, A.; Ecker, J.R.; Palm, C.J.; Federspiel, N.A.; Kaul, S.; White, O.;
Alonso, J.; Altaf, H.; Araujo, R.; Bowman, C.L.; Brooks, S.Y.; Buehler, E.;
Chan, A.; Chao, Q.; Chen, H.; Cheuk, R.F.; Chin, C.W.; Chung, M.K.; Conn, L.;
Conway, A.B.; Conway, A.R.; Creasy, T.H.; Dewar, K.; Dunn, P.; Etgu, P.;
Feldblyum, T.V.; Feng, J.; Fong, B.; Fujii, C.Y.; Gill, J.E.; Goldsmith, A.D.;
Haas, B.; Hansen, N.F.; Hughes, B.; Huizar, L.
Nature 408, 816-820, 2000
A; Authors: Hunter, J.L.; Jenkins, J.; Johnson-Hopson, C.; Khan, S.; Khaykin, E.;
Kim, C.J.; Koo, H.L.; Kremenetskaia, I.; Kurtz, D.B.; Kwan, A.; Lam, B.; Langin-
Hooper, S.; Lee, A.; Lee, J.M.; Lenz, C.A.; Li, J.H.; Li, Y.; Lin, X.; Liu,
S.K., Liu, E.A.; Luros, J.S.; Maiti, R.; Marziali, A.; Militscher, J.; Miranda,
M.: Nguyen, M.; Nierman, W.C.; Osborne, E.I.; Pai, G.; Peterson, J.; Pham, P.K.;
Rizzo, M.; Rooney, T.; Rowley, D.; Sakano, H.
A; Authors: Salzberg, S.L.; Schwartz, J.R.; Shinn, P.; Southwick, A.M.; Sun, H.;
Tallon, L.J.; Tambunga, G.; Toriumi, M.J.; Town, C.D.; Utterback, T.; van Aken,
S.; Vaysberg, M.; Vysotskaia, V.S.; Walker, M.; Wu, D.; Yu, G.; Fraser, C.M.;
Venter, J.C.; Davis, R.W.
A; Title: Sequence and analysis of chromosome 1 of the plant Arabidopsis.
A; Reference number: A86141; MUID:21016719; PMID:11130712
A; Accession: D86190
A; Status: preliminary
A; Molecule type: DNA
A; Residues: 1-1136 <STO>
A;Cross-references: UNIPROT:Q9SYK4; GB:AE005172; NID:q4836913; PIDN:AAD30615.1;
GSPDB.GN00141
C; Genetics:
A; Map position: 1
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                         15.3%; Score 309.5; DB 2; Length 1136;
                         27.3%; Pred. No. 4.3e-18;
 Best Local Similarity
 Matches
          93; Conservative 52; Mismatches 109; Indels
                                                              87; Gaps
            8 SFIKVSHVRMQGILLLVFAKYQHLPYIQILSTKSTPTGLFGYWGNKGGVNICLKLYGYYV 67
Qу
              : | :: |: |: |: |: | : | | : | |
                                                    651 TFERMGSRQLAGLLISLWARKDIRTHVGDLDVAAVPCGFGRAIGNKGGVGLRIRVYDRIM 710
Db
          68 SIINCHLPPHIS-------NYQRLEHFDRILEMQNC------EGRDIP--- 103
Qу
                                  |: || | : |
          711 CFVNCHLAAHLEAVNRRNADFNHIFRLMVFSRGQNLSNAAAAGVSTSAYTTKSNTIPSTG 770
ďŒ
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104 -----NILDHDLIIWFGDMNFRIEDFGLHF--VRESIKNRCYGGLWEKDQLSIAKKHDP 155
Qy
                        ]:: :||| |:|: ||: : |: | : | : | |:|||
         771 AEEIKSDLAAADMVAFFGDFNYRL--FGITYDEARDFISORSFDWLRERDQLRAEMKVGK 828
Db
         156 LLREFQEGRLLFPPTYKFDRNSN---DYDTSEKKRKPAWTDRILWRLKRQPCAGPDTP1P 212
Qу
                                        ||: |||| ||:::|
              829 VFQGMREALITFPPTYKFERNRSGLGGYDSGEKKRIPAWCDRVIYR------DTQSS 879
Db
         213 PASHFSL-----SLRGYSSHMTYGISDHKPVSGTF----- 242
Qу
                         | | :|
         880 PFSESNLOCPVVSSVIMYEACMDVTESDHKPVRCKFHATIAHVDKSVRROELGKIIRSNE 939
Db
         243 -----DLELKP--LVSAPLIVLMPED--LWTVENDMMVS 272
Qy
                    Db
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RESULT 8
T00670
probable inositol polyphosphate 5'-phosphatase [imported] - Arabidopsis thaliana
N; Alternate names: hypothetical protein F6E13.3
C; Species: Arabidopsis thaliana (mouse-ear cress)
C;Date: 01-Feb-1999 #sequence revision 01-Feb-1999 #text change 09-Jul-2004
C; Accession: T00670; H84871
R; Rounsley, S.D.; Kaul, S.; Lin, X.; Ketchum, K.A.; Crosby, M.L.; Brandon, R.C.;
Sykes, S.M.; Mason, T.M.; Kerlavage, A.R.; Adams, M.D.; Somerville, C.R.;
Venter, J.C.
submitted to the EMBL Data Library, June 1998
A; Description: Arabidopsis thaliana chromosome II BAC F6E13 genomic sequence.
A; Reference number: Z14180
A; Accession: T00670
A; Status: translated from GB/EMBL/DDBJ
A; Molecule type: DNA
A; Residues: 1-1305 < ROU>
A; Cross-references: UNIPROT: 080560; EMBL: AC004005; NID: q3212846; PID: q3212848
A; Experimental source: cultivar Columbia
R; Lin, X.; Kaul, S.; Rounsley, S.D.; Shea, T.P.; Benito, M.I.; Town, C.D.;
Fujii, C.Y.; Mason, T.M.; Bowman, C.L.; Barnstead, M.E.; Feldblyum, T.V.; Buell,
C.R.; Ketchum, K.A.; Lee, J.J.; Ronning, C.M.; Koo, H.; Moffat, K.S.; Cronin,
L.A.; Shen, M.; VanAken, S.E.; Umayam, L.; Tallon, L.J.; Gill, J.E.; Adams,
M.D.; Carrera, A.J.; Creasy, T.H.; Goodman, H.M.; Somerville, C.R.; Copenhaver,
G.P.; Preuss, D.; Nierman, W.C.; White, O.; Eisen, J.A.; Salzberg, S.L.; Fraser,
C.M.; Venter, J.C.
Nature 402, 761-768, 1999
A; Title: Sequence and analysis of chromosome 2 of the plant Arabidopsis
A; Reference number: A84420; MUID: 20083487; PMID: 10617197
A; Accession: H84871
A; Status: preliminary
A; Molecule type: DNA
A; Residues: 1-1305 <STO>
A; Cross-references: GB: AE002093; NID: g3212848; PIDN: AAC23399.1; GSPDB: GN00139
C; Genetics:
A;Gene: F6E13.3; At2g43900
A; Map position: 2
A; Introns: 278/1; 362/2; 631/3; 674/2; 702/3; 758/1; 771/3; 861/1; 902/2; 1022/3
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15.1%; Score 306; DB 2; Length 1305;
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 Best Local Similarity 27.1%; Pred. No. 1e-17;
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          9 FIKVSHVRMQGILLLVFAKYQHLPYIQILSTKSTPTGLFGYWGNKGGVNICLKLYGYYVS 68
Qy
                  658 FERMGSROLAGLLISLWVRKNLRTHVGDIDVAAVPCGFGRAIGNKGGVGLRIRVFDRIMC 717
Db
         69 IINCHLPPHISNNYORLEHFDRILE----MQNCEGRDIP------ 103
Qу
             718 FINCHLAAHLEAVNRRNADFDHIYKTMSFTRSSNAHNAPAAGVSTGSHTTKSANNANVNT 777
Db
        104 ----NILDHDLIIWFGDMNFRIEDFGLHF--VRESIKNRCYGGLWEKDQLSIAKKHDPL 156
QУ
                778 EETKODLAEADMVVFFGDFNYRL--FGISYDEARDFVSQRSFDWLREKDQLRAEMKAGRV 835
Db
        157 LREFQEGRLLFPPTYKFDRNS---NDYDTSEKKRKPAWTDRILWRLKRQPCAGPDTPIPP 213
Qу
             836 FQGMREAIITFPPTYKFERHRPGLGGYDSGEKKRIPAWCDRVIFR-----DTRTSP 886
Db
        214 ASHFSL-----SLRGYSSHMTYGISDHKPVSGTFDLELK 247
Qу
                     887 ESECSLDCPVVASIMLYDACMDVTESDHKPVRCKFHVKIE 926
Db
RESULT 9
T19338
hypothetical protein C16C2.3 - Caenorhabditis elegans
C; Species: Caenorhabditis elegans
C;Data: 15-Oct-1999 #sequence_revision 15-Oct-1999 #text_change 09-Jul-2004
C; Accession: T19338
R; Barlow, K.
submitted to the EMBL Data Library, October 1996
A; Reference number: Z19109
A; Accession: T19338
A; Status: preliminary; translated from GB/EMBL/DDBJ
A; Molecule type: DNA
A; Residues: 1-753 <WIL>
A; Cross-references: UNIPROT:017590; EMBL:Z81036; PIDN:CAB02743.1; GSPDB:GN00019;
CESP:C16C2.3
A; Experimental source: clone C16C2
C:Genetics:
A; Gene: CESP: C16C2.3
A; Mar position: 1
A; Introns: 17/1; 36/2; 88/2; 131/2; 170/3; 268/3; 389/2; 475/3; 498/3; 632/1;
677/2; 713/1
                       14.9%; Score 300.5; DB 2; Length 753;
  Query Match
  Best Local Similarity 31.1%; Pred. No. 1.4e-17;
         85; Conservative 40; Mismatches 93; Indels 55; Gaps 9;
 Matches
           9 FIKVSHVRMQGILLLVFAKYQHLPYIQILSTKSTPTG---LFGYWGNKGGVNICLKLYGY 65
Qу
                               : :: | || | | | || || || : :|:
            | | :|: || ::||
         168 FNVVGSMRLVGIFVIVFQAVNSKVRVSDVNVKYVATGISVLVNKLGNKGGTAVSMKMNDT 227
Db
          66 YVSIINCHLPPHISNNYQRLEHFDRILEMQNCEGRDIPN------ 104
Ov
            : | : | | | |
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Db
         228 WVCFVNAHFA--AGNNE------LERRNQDFRDIYNDMVFYPRSQQEGLRDRPLEVP 276
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Qу
                : : | : | |
         277 IMCLYDHDVVFWFGDLNYRLNTDMYGISNDEVRRIASSDKFADLLQHCQLREQMARGTVF 336
Db
         158 REFQE-GRLLFPPTYKFDRNSNDYDTSEKKRKPAWTDRILWRLKRQPCAGPDTPIPPASH 216
Qу
             337 KDFEEPSTLPFRPTYKYDCGTNTWDTSEKGRVPAWTDRIL-TFKKYPQVGLE----- 387
Db
         217 FSLSLRGYSSHMTYGISDHKPVSGTFDLELKPL 249
QУ
                1:1 | | | | | | | | | | | | | | | | |
         388 ---SIRPMVSIDTITISDHKPVRAMFNLKVKKI 417
Db
RESULT 10
D96515
hypothetical protein F16N3.22 [imported] - Arabidopsis thaliana
C; Species: Arabidopsis thaliana (mouse-ear cress)
C;Date: 02-Mar-2001 #sequence_revision 02-Mar-2001 #text_change 09-Jul-2004
C; Accession: D96515
R; Theologis, A.; Ecker, J.R.; Palm, C.J.; Federspiel, N.A.; Kaul, S.; White, O.;
Alonso, J.; Altaf, H.; Araujo, R.; Bowman, C.L.; Brooks, S.Y.; Buehler, E.;
Chan, A.; Chao, Q.; Chen, H.; Cheuk, R.F.; Chin, C.W.; Chung, M.K.; Conn, L.;
Conway, A.D.; Conway, A.R.; Creasy, T.H.; Dewar, K.; Dunn, P.; Etgu, P.;
Feldblyum, T.V.; Feng, J.: Fong, B.; Fujii, C.Y.; Gill, J.E.; Goldsmith, A.D.;
Haas, B.; Hansen, N.F.; Hughes, B.; Huizar, L.
Nature 403, 816-820, 2000
A; Authors: Hunter, J.L.; Jenkins, J.; Johnson-Hopson, C.; Khan, S.; Khaykin, E.;
Kim. C.J.; Koo, H.L.; Kremenetskaia, I.; Kurtz, D.B.; Kwan, A.; Lam, B.; Langin-
Hooper, S.; Lee, A.; Lee, J.M.; Lenz, C.A.; Li, J.H.; Li, Y.; Lin, X.; Liu,
S.X.; Liu, Z.A.; Luros, J.S.; Maiti, R.; Marziali, A.; Militscher, J.; Miranda,
M.; Mguyen. M.; Nierman, W.C.; Osborne, B.I.; Pai, G.; Peterson, J.; Pham, P.K.;
Rizzo, M.; Rooney, T.; Rowley, D.; Sakano. H.
A; Authors: Salzberg, S.L.; Schwartz, J.R.; Shinn, P.; Southwick, A.M.; Sun, H:;
Tallon, L.J.; Tambunga, G.; Toriumi, M.J.; Town, C.D.; Utterback, T.; van Aken,
S.; Vaysberg, M.; Vysotskaia, V.S.; Walker, M.; Wu, D.; Yu, G.; Fraser, C.M.;
Venter, J.C.; Davis, R.W.
A; Title: Sequence and analysis of chromosome 1 of the plant Arabidopsis.
A; Reference number: A86141; MUID: 21016719; PMID: 11130712
A; Accession: D96515
A; Status: preliminary
A; Molecule type: DNA
A; Residues: 1-331 <STO>
A;Cross-references: UNIPROT:Q9SX81; GB:AE005173; NID:g5668810; PIDN:AAD46036.1;
GSPDB:GN00141
C; Genetics:
A; Gene: F16N3.22
A; Map position: 1
                         14.8%; Score 298.5; DB 2; Length 331;
  Query Match
                        34.4%; Pred. No. 6.8e-18;
  Best Local Similarity
          83; Conservative 33; Mismatches
                                               90; Indels
                                                             35; Gaps
                                                                         9;
          16 RMQGILLLVFA-KYQHLPYIQILSTKSTPTGLFGYWG-NKGGVNICLKLYGYYVSIINCH 73
Qу
             Db
         113 KLQSVQLYLFGPKNSHTLVKELKAERYSVGGCGGLIGRKKGAVAIRINYDDIKMVFISCH 172
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74 LPPHISNNYQRLEHFDRILEMONCEGRDIPNIL-----DHDLIIWFGDMNFRIEDFGLH 127
Qу
                            :: :| | | | | |
                                                 173 LSAHAKK------VDQRNTELRHIANSLLPRDKRKRDLTVWLGDLNYRIQDVSNH 221
Db
         128 FVRESIKNRCYGGLWEKDQLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDYDTSEKKR 187
QУ
              222 PVRSLIONHLOSVLVSKDOLLOEAERGEIFKGYSEGTLGFKPTYKYNVGSSDYDTSHKIR 281
Db
         188 KPAWTDRILWRLKROPCAGPDTPIPPASHFSLSLRGYSS-HMTYGISDHKPVSGTFDLEL 246
Qу
                                  11
         282 VPAWTDRILFKIQ------DT-----DNIQATLHSYDSIDQVYG-SDHKPVKA--DLCL 326
Db
Qу
         247 K 247
         327 K 327
Dh
RESULT 11
G84792
hypothetical protein At2g37440 [imported] - Arabidopsis thaliana
C; Species: Arabidopsis thaliana (mouse-ear cress)
C;Date: 02-Feb-2001 #sequence_revision 02-Feb-2001 #text change 09-Jul-2004
C; Accession: G84792
R; Lin, X.; Kaul, S.; Rounsley, S.D.; Shea, T.P.; Benito, M.I.; Town, C.D.;
Fujii, C.Y.; Mason, T.M.; Bowman, C.L.; Barnstead, M.E.; Feldblyum, T.V.; Buell,
C.R.; Ketchum, K.A.; Lee, J.J.; Ronning, C.M.; Koo, H.; Moffat, K.S.; Cronin,
L.A.; Shen, M.; VanAken, S.E.; Umayam, L.; Tallon, L.J.; Gill, J.E.; Adams,
M.D.; Carrera, A.J.; Creasy, T.H.; Goodman, H.M.; Somerville, C.R.; Copenhaver,
G.P.; Preuss, D.; Nierman, W.C.; White, O.; Eisen, J.A.; Salzberg, S.L.; Fraser,
C.M.; Venter, J.C.
Nature 402, 761-768, 1999
A; Title: Sequence and analysis of chromosome 2 of the plant Arabidopsis
thaliana.
A; Reference number: A84420; MUID: 20083487; PMID: 10617197
A; Accession: G84792
A; Status: preliminary
A; Molecule type: DNA
A; Residues: 1-401 <STO>
A; Cross-references: UNIPROT: Q9ZUS3; GB: AE002093; NID: q4056496; PIDN: AAC98062.1;
GSPDB:GN00139
C; Genetics:
A; Gene: At2q37440
A; Map position: 2
                       14.7%; Score 297.5; DB 2; Length 401;
 Query Match
 Best Local Similarity 29.7%; Pred. No. 1.1e-17;
          76; Conservative 43; Mismatches 98; Indels
                                                          39; Gaps 7;
 Matches
           6 PLSFIKVSHVRMQGILLLVFAKYQHLPYIQILSTKSTPTGLFGYWGNKGGVNICLKLYGY 65
Qу
             113 PRGYSLAASKOMVGIFLCVWVRDDLRKRITNLKVSCVGRGIMGYLGNKGSVSISMSLHET 172
Db
          66 YVSIINCHLP-----PHISNNYQRLEHFDRILEMQNCEGRDIPNILDHDLIIWFGDMNF 119
Qу
                            : | | | :::
                                                      |:||| :|| ||:|:
              : : ||
ď.Q
         173 SLCFVCTHLTSGEKEGDELRRNLDVTEIFKRTRFSRSSKDSRPETIMDHDKVIWLGDLNY 232
```

```
120 RIE-DFGLHFVRESIKNRCYGGLWEKDOLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSN 178
Qу
                       233 RLRASSDLH---EQLRNHDWESLLEKDQLKIEQRAGRIFKGWEEGKIYFAPTYKYRINSD 289
Db
         179 DY-----DTSEKKRKPAWTDRILWR---LKRQPCAGPDTPIPPASHFSLSLRGYSSHMTY 230
Qу
             290 NYVVQTEKSKEKRRTPAWCDRILWKGDGMKQ------LWYVRGESK---- 329
Db
         231 GISDHKPVSGTFDLEL 246
Qу
              Dh
         330 -FSDHRPVOSLFSVHI 344
RESULT 12
S61667
probable membrane protein YOR109w - yeast (Saccharomyces cerevisiae)
N; Alternate names: hypothetical protein 03231; hypothetical protein YOR3231w
C; Species: Saccharomyces cerevisiae
C;Date: 09-Mar-1996 #sequence_revision 12-Apr-1996 #text_change 09-Jul-2004
C; Accession: S61667; S66994
R; Benes, V.; Andrade, M.A.; Rechmann, S.; Teodoru, C.; Banrevi, A.; Sander, C.;
Valencia, A.; Ansorge, W.; Voss, H.
submitted to the EMBL Data Library, December 1995
A; Description: Nucleotide sequence and analysis of a 130 kb fragment of yeast
chromosome XV.
A; Reference number: S61643
A; Accession: S61667
A; Molecule type: DNA
A; Residues: 1-1107 <BEN>
A;Cmoss-references: UNIPROT:Q12271; EMBL:X94335; NID:g1262139; PID:g1164954
R; Voss, H.; Benes, V.; Rechmann, S.; Teodoru, C.; Schwager, C.; Paces, V.;
Ansorge, W.
submitted to the Protein Sequence Database, July 1996
A;Reference number: S66965
A; Accession: S66994
A; Molecule type: DNA
A; Residues: 1-1107 < VOS>
A;Cross-references: EMBL:Z75017; NID:g1420294; PID:g1420295; MIPS:YOR109w
A; Experimental source: strain S288C
C; Genetics:
A; Gene: SGD: INP53
A; Cross-references: SGD:S0005635; MIPS:YOR109w
A; Map position: 15R
C; Keywords: transmembrane protein
F;60-76/Domain: transmembrane #status predicted <TM1>
F;642-658/Domain: transmembrane #status predicted <TM2>
 Query Match
                        14.4%; Score 291.5; DB 2; Length 1107;
 Best Local Similarity
                       26.8%; Pred. No. 1.5e-16;
         98; Conservative 57; Mismatches 141; Indels
 Matches
                                                           69; Gaps
          16 RMQGILLLVFAKYQHLPYIQILSTKSTPTGLFGYWGNKGGVNICLKLYGYYVSIINCHLP 75
Qу
                         : | : | : | |
         648 QMTSLLILFFVKADKAKYVKQVEGATKKTGFRGMAGNKGAVSIRFEYGATSFCFVNSHLA 707
Db
          76 PHISNNYQRLEHFDRILE-MONCEGRDIPNILDHDLIIWFGDMNFRI----EDFGLHFVR 130
Qy
                :| :| :: |: : : ||: || | | | |||:||
```

```
708 AGATNVEERRSDYESIVRGITFTRTKMIPH---HDSIFWLGDMNYRINLPNED-----VR 759
Db
         131 ESIKNRCYG---GLWEKDQLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDYDTSEKKR 187
Qу
               760 RELLNOEEGYIDKLLHFDOLTLGINSGSVFEGFKEPTLKFRPTYKYDPGTGTYDSSEKER 819
Db
         188 KPAWTDRILWRLKR-QPCAGPDTPIPPASHFSLSLRGYSSHMTYGISDHKPVSGTF---- 242
QУ
              1:||||:::: | | : | ||
         820 TPSWTDRIIYKGENLLPLSYSDAPIM------ISDHRPVYAAYRAKI 860
Db
Qу
         243 -----DLELKPLVSAPLIVLMPEDLWTVENDMM---VSYSSTSDFPSSPWDWIGLYKV 292
                   861 TFVDDKERLSLKKRLFTEYKQEHPEEPGSLISDLLSLDLDNKSTDGFKSSS-----ES 913
Db
Qy
         293 GLRDVNDYVSYAWVGDSKVSCSDNLNOVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRP 352
              914 SLLDIDPIMAQPTA--SSVASSSPVSSASASLQPVRT-----QNSSQSRTPIKKP 961
Db
         353 FOIPP 357
ŌУ
               Db
         962 VLRPP 966
RESULT 13
548433
inositol-1,4,5-trisphosphate 5-phosphatase homolog YIL002c - yeast
(Saccharomyces cerevisiae)
N; Alternate names: protein YIA2c
C; Species: Saccharomyces cerevisiae
C;Date: 02-Dec-1994 #sequence revision 02-Dec-1994 #text change 09-Jul-2004
C; Accession: S48433; S58703; S50873
R; Badcock, K.; Churcher, C.
submitted to the EMBL Data Library, August 1994
A; Reference number: S48432
A; Accession: S48433
A; Molecule type: DNA
A; Residues: 1-946 <BAD>
A; Cross-references: UNIPROT: P40559; GB: Z47047; EMBL: Z38062; NID: g603997;
PID:g763344; MIPS:YIL002c
R; Voss, H.
submitted to the EMBL Data Library, June 1994
A; Reference number: S58703
A; Accession: S53703
A; Molecule type: DNA
A; Residues: 1-946 < VOS>
A; Cross-references: EMBL: X79743
R; Voss, H.; Tamames, J.; Teodoru, C.; Valencia, A.; Sensen, C.; Wiemann, S.;
Schwager, C.; Zimmermann, J.; Sander, C.; Ansorge, W.
Yeast 11, 61-78, 1995
A;Title: Nucleotide sequence and analysis of the centromeric region of yeast
chromosome IX.
A; Reference number: S50795; MUID: 95282515; PMID: 7762303
A: Accession: S50873
A; Status: nucleic acid sequence not shown
A; Molecule type: DNA
A; Residues: 650-820 < VOW>
A; Cross-references: EMBL: X79743
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```
C; Genetics:
A; Gene: SGD: INP51
A; Cross-references: SGD:S0001264; MIPS:YIL002c
A; Map position: 9L
C; Keywords: transmembrane protein
F;71-87/Domain: transmembrane #status predicted <TMM>
                       14.3%; Score 289.5; DB 2; Length 946;
 Query Match
 Best Local Similarity 31.5%; Pred. No. 1.7e-16;
         90; Conservative 38; Mismatches 115; Indels
                                                          43;
                                                              Gaps
                                                                      8;
           9 FIKVSHVRMOGILLLVFAKYOHLPYIQILSTKSTPTGLFGYWGNKGGVNICLKLYGYYVS 68
Qy
                 :: |||||:|
                                   ::::
                                            |\cdot|\cdot|\cdot|\cdot|
         607 YIRLWSTQLGGILLLLFMNETEYSKVKHIEGDVKKTGFGGMASNKGAVAVSFKYSATRFC 666
Db
          69 IINCHLPPHISNNYORLEHFDRILE-MONCEGRDIPNILDHDLIIWFGDMNFRI----ED 123
Qу
             :: || : | : : : :
                                            667 VLVSHLAAGLENVEQRHNDYKTIAKSIRFSKGL---RIKDHDAIIWMGDFNYRILMSNED 723
Db
         124 FGLHFVRESIKNRCYGGLWEKDQLS---IAKKHDPLLREFQEGRLLFPPTYKFDRNSNDY 180
Qу
                 724 ----VRRKIVSKEYASLFEKDQLNQQMIAGESFPY---FHEMAIDFPPTYKFDPGTKNY 775
Db
         181 DTSEKKRKPAWTDRILWRLKRQPCAGPDTPIPPASHFSLSLRGYSSHMTYGISDHKPVSG 240
Qу
             Db
         241 TF-----DLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDF 279
Q٧
                    819 IFRARVTVVDEQKKTTLGTQIYEKIMERLEGLDDDEKIAVLSDDAF 864
DΡ
RESULT 14
F84725
probable inositol polyphosphate 5'-phosphatase [imported] - Arabidopsis thaliana
C; Species: Arabidopsis thaliana (mouse-ear cress)
C;Date: 02-Feb-2001 #sequence revision 02-Feb-2001 #text_change 09-Jul-2004
C; Accession. F84725
R; Lin, X.; Kaul, S.; Rounsley, S.D.; Shea, T.P.; Benito, M.I.; Town, C.D.;
Fujii, C.Y.; Mason, T.M.; Bowman, C.L.; Barnstead, M.E.; Feldblyum, T.V.; Buell,
C.R.; Ketchum, K.A.; Lee, J.J.; Ronning, C.M.; Koo, H.; Moffat, K.S.; Cronin,
L.A.; Shen, M.; VanAken, S.E.; Umayam, L.; Tallon, L.J.; Gill, J.E.; Adams,
M.D.; Carrera, A.J.; Creasy, T.H.; Goodman, H.M.; Somerville, C.R.; Copenhaver,
G.P.; Preuss, D.; Nierman, W.C.; White, O.; Eisen, J.A.; Salzberg, S.L.; Fraser,
C.M.; Venter, J.C.
Nature 402, 761-768, 1999
A; Title: Sequence and analysis of chromosome 2 of the plant Arabidopsis
A; Reference number: A84420; MUID: 20083487; PMID: 10617197
A; Accession: F84725
A; Status: preliminary
A; Molecule type: DNA
A; Residues: 1-1144 <STO>
A;Cross-references: UNIPROT:Q9SKB7; GB:AE002093; NID:g4887753; PIDN:AAD32289.1;
GSPDB:GN00139
C; Genetics:
A; Gene: At2g31830
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Query Match
                        14.1%; Score 284; DB 2; Length 1144;
 Best Local Similarity 27.4%; Pred. No. 6.8e-16;
 Matches
           75; Conservative 49; Mismatches
                                             94; Indels
                                                           56; Gaps
           8 SFIKVSHVRMQGILLLVFAKYQHLPYIQILSTKSTPTGLFGYWGNKGGVNICLKLYGYYV 67
Qу
             :| :: :: |:|: :: :: | : | |
                                                     111111 : :::1
Db
         660 TFERMGSRQLAGLLISLWVRKSIRTHVGDLDVAAVPCGFGRAIGNKGGVGLRIRVYDRIM 719
          68 SIINCHLPPHISNNYQRLEHFDRIL-EMQNCEGRDI------PN---- 104
QУ
               720 CFVNCHLAAHLEAVTRRNADFNHIYRSMVFSKGOSVYTAAAAGASTSAOALKNNPNTNNS 779
Db
         105 -----ILDHDLIIWFGDMNFRIEDFGLHF--VRESIKNRCYGGLWEKDQLSIAKKHDP 155
QУ
                       730 TEEEKSHLASADLVAFFGDFNYRL--FGITYDEARDFISHRSFDWLREKDQLRQEMNEGK 837
Db
         156 LLREFQEGRLLFPPTYKFDRNS---NDYDTSEKKRKPAWTDRILWRLKRQPCAGPDTPIP 212
Qу
             838 VFQGMREALITFPPTYKFEKNKPGLGGYDSGEKKRIPAWCDRVIYRDNQS----- 887
Db
         213 PASHFSLSLR------GYSSHMTYGISDHKPV 238
Qу
               |: ||:
                              | : |
         888 -ISYTECSLKCPVVSSTIMYEACMDVTESDHKPV 920
dG
RESULT 15
JC5765
inositol polyphosphate 5-phosphatase (EC 3.1.3.-) 2 - human
C; Species: Momo sapiens (man)
C;Date 24-Jan-1998 #sequence revision 13-Mar-1998 #text change 09-Jul-2004
C; Accession: JC5765
R; Pesesse, X.; Deleu, S.; De Smedt, F.; Drayer, L.; Erneux, C.
Biochem. Biophys. Res. Commun. 239, 697-700, 1997
A; Title: Identification of a second SH2-domain-containing protein closely
related to the phosphatidylinositol polyphosphate 5-phosphatase SHIP.
A;Reference number: JC5765; MUID:98042455; PMID:9367831
A; Accession: JC5765
A; Status: nucleic acid sequence not shown
A; Molecule type: mRNA
A; Residues: 1-1258 < PES>
A; Cross-references: UNIPROT: 015357; GB: Y14385; NID: q2653423; PIDN: CAA74743.1;
PID:c1108709; PID:g2653424
C; Comment: This enzyme plays a role in metabolism of inositol phosphate and
phosphatidylinositol.
C; Keywords: phosphoric monoester hydrolase
F;21-117/Domain: SH2 homology <SH2>
F;427-729/Domain: catalytic #status predicted <CAT>
F;927-1171/Region: proline-rich
                               Score 284; DB 2; Length 1258;
  Query Match
                        14.1%;
                        25.0%; Pred. No. 7.7e-16;
  Best Local Similarity
  Matches 96; Conservative 66; Mismatches 154; Indels
                                                           68; Gaps
           4 LSPLSFIKVSHVRMQGILLLVFAKYQHLPYIQILSTKSTPTGLFGYWGNKGGVNICLKLY 63
Qу
```

A; Map position: 2

Db	492	LTDLDYRPIAMQSLWNIKVAVLVKPEHENRISHVSTSSVKTGIANTLGNKGAVGVSFMF							
Qу	64	GYYVSIINCHLPPHISNNYQRLEHFDRILEMQNCEGRDIPNILDHDLIIWFGDM							
Db	552	GTSFGFVNCHLTSGNEKTARRNQNYLDILRLLSLGDRQL-NAFDISLRFTHLFWFGDLN							
Qу	120	RIEDFGLHFVRESIKNRCYGGLWEKDQLSIAK-KHDPLLREFQEGRLLFPPTYKFDRNSN : : : : :	178						
Db	611	: : : : : ::: : : ::: RL-DMDIQEILNYISRKEFEPLLRVDQLNLEREKHKVFLR-FSEEEISFPPTYRYERGS							
Qу		DYDTSEKKRKPAWTDRILWRLKRQPCAGPDTPIPPASHFSLSLRGYSSHMTYC							
Db	669								
QУ	232	ISDHKPVSGTFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDW1							
ďū	715	TSDHSPVFGTFEVGVTSQFISKKGLSKTSDQAYIEFESIEAIVKTASRTKFFIE	768						
Qy	289	LYKVGLRDVNDYVSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLL							
Db	769	: :: : : : : FYSTCLEEYKKSFENDAQSSDNINFLKVQWSSRQLPTLKPILADIEYLQDQHLL 322							
Qу	337	CYYSNSLRSVVG 348 : : : ::							
œ	323	: : : :: LTVKSMDGYESYGECVVALKSMIG 846							
Search sompleted: February 10, 2005, 17:50:42 Job time: 44 secs									
		GenCore version 5.1.6 Copyright (c) 1993 - 2005 Compugen Ltd.							
OM protein - protein search, using sw model									
Run on:		February 10, 2005, 17:38:56 ; Search time 176 Seconds (without alignments) 1082.349 Million cell updates/se	ec						
Perfect score: Sequence:									
Scoring table:		BLOSUM62 Gapop 10.0 , Gapext 0.5							
Searched:		1612378 seqs, 512079187 residues							
Total number of hits satisfying chosen parameters: 1612378									
Minimum DB seq length: 0 Maximum DB seq length: 2000000000									
Post-processing: Minimum Match 0%									

Maximum Match 100%

Listing first 45 summaries

Database :

39

40

41

42

43

327.5

327.5

326

326

326

UniProt 03:*

1: uniprot_sprot:*
2: uniprot_trembl:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

					SUMMARIES	
		€				
Result		Query				
No.	Score	Match	Length	DB	ID	Description
1	2010	99.5	448	1	SKIP_HUMAN	Q9bt40 homo sapien
2	1525	75.5	468	1	SKIP MOUSE	Q8c5l6 mus musculu
3	1115	55.2	496	2	Q6GN99	Q6gn99 xenopus lae
4	873.5	43.2	1003	1	P5PA_MOUSE	P59644 mus musculu
5	872.5	43.2	1001	1	P5PA_RAT	Q9jmc1 rattus norv
6	870.5	43.1	638	2	Q6ZTH6	Q6zth6 homo sapien
. 7	864.5 -	42.8	1006	1	P5PA_HUMAN	Q15735 homo sapien
8	516	25.5	501	2	Q7PYY6	Q7pyy6 anopheles g
9	507.5	25.1	508	2	Q9VXE7	Q9vxe7 drosophila
10	460.5	22.8	357	2	Q9V7X0	Q9v7x0 drosophila
11	432.5	21.4	736	2	Q86YE1	Q86yel homo sapien
12	432.5	21.4	748	2	Q6PD53	Q6pd53 homo sapien
1.3	432.5	21.4	871	2	Q6P6D4	Q6p6d4 homo sapien
14	432.5	21.4	942	1	I5P2_HUMAN	P32019 homo sapien
15	423.5	21.0	742	2	Q8CF65	Q8cf65 mus musculu
16	423.5	21.0	757	2	Q9DBW2	Q9dbw2 mus musculu
17	423.5	21.0	776	2	054996	054996 mus musculu
18	423.5	21.0	825	2	Q91ZF8	Q91zf8 mus musculu
19	423.5	21.0	993	2	Q8K337	Q8k337 mus musculu
20	334	19.0	787	2	Q8I7P3	Q8i7p3 dictyosteli
21	373.5	18.5	727	2	Q8BXC9	Q8bxc9 mus musculu
22	373.5	18.5	900	2	Q6NVF0	Q6nvf0 mus musculu
23	372	18.4	511	2	Q8BXT3	Q8bxt3 mus musculu
24	367.5	18.2	901	1	OCRL_HUMAN	Q01968 homo sapien
25	341.5	16.9	1800	2	Q8I7P5	Q8i7p5 dictyosteli
26	340	16.8	1113	2	Q9XUD3	Q9xud3 caenorhabdi
27	340	16.8	1119	2	Q9GT42	Q9gt42 caenorhabdi
28	339		1101	2	Q84W55	Q84w55 arabidopsis
. 29	339	16.8	1291	2	Q9DGN7	Q9dgn7 lampetra fl
30 .	338	16.7	1101	2	Q9XFT1	Q9xft1 arabidopsis
31		16.6	1273	2	Q7Q001	Q7q001 anopheles g
32	334	16.5	1434	1	SYJ2_MOUSE	Q9d2g5 mus musculu
33	328	16.2	552	2	Q6AVN0	Q6avn0 oryza sativ
34	328	16.2	1315	2	094984	094984 homo sapien
35	328	16.2	1574	1	SYJ1_MOUSE	Q8chc4 mus musculu
36	328	16.2	1574	1	SYJ1_RAT	Q62910 rattus norv
37	328	16.2	1575	1	SYJ1_HUMAN	043426 homo sapien
38	327.5	16.2	1288	2	Q8IZA8	Q8iza8 homo sapien

015056 homo sapien

Q9h226 homo sapien O55207 rattus norv

Q91zd9 rattus norv

Q91zd8 rattus norv

16.2 1443 1 SYJ2_HUMAN

16.2 1496 2 Q9H226

16.1 1451 2 Q91ZD9

16.1 1496 2 Q91ZD8

16.1 1248 1 SYJ2_RAT

ALIGNMENTS

```
RESULT 1
SKIP HUMAN
     SKIP HUMAN
                    STANDARD;
                                   PRT:
                                          448 AA.
AC
     Q9BT40; Q15733; Q9NPJ5; Q9P2R5;
DT
     10-OCT-2003 (Rel. 42, Created)
DΤ
     10-OCT-2003 (Rel. 42, Last sequence update)
DT'
     05-JUL-2004 (Rel. 44, Last annotation update)
DE
     Skeletal muscle and kidney enriched incsitol phosphatase
DE
     (EC 3.1.3.56).
GN
     Name=SKIP; Synonyms=PPS;
os
     Homo sapiens (Human).
OC
     Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
     Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
OX
     NCBI TaxID=9606;
RN
RΡ
     SEQUENCE FROM N.A. (ISOFORMS 1 AND 2), FUNCTION, AND TISSUE
RΡ
     SPECIFICITY.
RC
     TISSUE=Testis:
     MEDLINE=20219123; PubMed=10753883; DOI=10.1074/jbc.275.15.10870;
RХ
     Ijuin T., Mochizuki Y., Fukami K., Funaki M., Asano T., Takenawa T.;
RÆ.
RT
     "Identification and characterization of a novel inositol polyphosphate
RT
     5-phosphatase.";
     J. Biol. Chem. 275:10870-10875(2000).
RI.
RN
     SEQUENCE FROM N.A. (ISOFORM 1).
RΡ
RC
     TISSUE=Eye;
     MEDLINE=22388257; PubMed=12477932; DOI=10.1073/pnas.242603899;
RX
RA
     Strausberg R.L., Feingold E.A., Grouse L.H., Derge J.G.,
RA
     Klausner R.D., Collins F.S., Wagner L., Shenmen C.M., Schuler G.D.,
     Altschul S.F., Zeeberg B., Buetow K.H., Schaefer C.F., Bhat N.K.,
RA
     Hopkins R.F., Jordan H., Moore T., Max S.I., Wang J., Hsieh F.,
RA
     Diatchenko L., Marusina K., Farmer A.A., Rubin G.M., Hong L.,
RA
     Stapleton M., Soares M.B., Bonaldo M.F., Casavant T.L., Scheetz T.E.,
RA
R.۸
     Brownstein M.J., Usdin T.B., Toshiyuki S., Carninci P., Prange C.,
RΑ
     Raha S.S., Loquellano N.A., Peters G.J., Abramson R.D., Mullahy S.J.,
RA
     Posak S.A., McEwan P.J., McKernan K.J., Malek J.A., Gunaratne P.H.,
     Richards S., Worley K.C., Hale S., Garcia A.M., Gay L.J., Hulyk S.W.,
RA
     Villalon D.K., Muzny D.M., Sodergren E.J., Lu X., Gibbs R.A.,
R\Lambda
RA
     Fahey J., Helton E., Ketteman M., Madan A., Rodrigues S., Sanchez A.,
     Whiting M., Madan A., Young A.C., Shevchenko Y., Bouffard G.G.,
RA
     Blakesley R.W., Touchman J.W., Green E.D., Dickson M.C.,
RA
     Rodriguez A.C., Grimwood J., Schmutz J., Myers R.M.,
RA
     Butterfield Y.S.N., Krzywinski M.I., Skalska U., Smailus D.E.,
RA.
     Schnerch A., Schein J.E., Jones S.J.M., Marra M.A.;
RA
RT
     "Generation and initial analysis of more than 15,000 full-length human
RT
     and mouse cDNA sequences.";
RL
     Proc. Natl. Acad. Sci. U.S.A. 99:16899-16903(2002).
RN
     SEQUENCE OF 120-448 FROM N.A.
RP
RC
     TISSUE=Brain;
```

```
RA
     Nussbaum R.L.;
     Submitted (JAN-1996) to the EMBL/GenBank/DDBJ databases.
RI_{i}
RN
     SUBCELLULAR LOCATION, AND MUTAGENESIS OF TYR-349; ASP-361; TRP-362 AND
RP
RP
     MEDLINE=22538472; PubMed=12536145; DOI=10.1074/jbc.M209991200;
RX
     Gurung R., Tan A., Ooms L.M., McGrath M.J., Huysmans R.D.,
RA
     Munday A.D., Prescott M., Whisstock J.C., Mitchell C.A.;
RA
     "Identification of a novel domain in two mammalian inositol-
RT
     polyphosphate 5-phosphatases that mediates membrane ruffle
RT
     localization. The inositol 5-phosphatase SKIP localizes to the
RT
     endoplasmic reticulum and translocates to membrane ruffles following
RT
     epidermal growth factor stimulation.";
RT
RL
     J. Biol. Chem. 278:11376-11385(2003).
CC
     -!- FUNCTION: Inositol 5-phosphatase which acts on inositol 1,4,5-
CC
         trisphosphate, inositol 1,3,4,5-tetrakisphosphate,
         phosphatidylinositol 4,5-bisphosphate and phosphatidylinositol
CC
         3,4,5-triphosphate. Has 6-fold higher affinity for
CC
         phosphatidylinositol 4,5-bisphosphate than for inositol 1,4,5-
CC
         trisphosphate. May negatively regulate assembly of the actin
CC
CC
         cytoskeleton.
CC
     -!- CATALYTIC ACTIVITY: D-myo-inositol 1,4,5-trisphosphate + H(2)0 =
CC
        myo-inositol 1,4-bisphosphate + phosphate.
     -!- SUBCELLULAR LOCATION: Endoplasmic reticulum. Following stimulation
CC
CC
         with EGF, translocates to membrane ruffles.
CC
     -!- ALTERNATIVE PRODUCTS:
        Event=Alternative splicing; Named isoforms=2;
CC
CC
           IsoId=Q9BT40-1; Sequence=Displayed;
CC
CC
         Name=2;
CC
           IsoId=Q9BT40-2; Sequence=VSP 050612;
CC
     -!- TISSUE SPECIFICITY: Ubiquitously expressed with highest levels in
CC
         skeletal muscle, heart and kidney.
     -!- SIMILARITY: Belongs to the inositol-1,4,5-trisphosphate 5-
CC
CC
         phosphatase type II family.
        ______
CC
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CC
     between the Swiss Institute of Bioinformatics and the EMBL outstation -
CC
CC
     the European Bioinformatics Institute. There are no restrictions on its
     use by non-profit institutions as long as its content is in no way
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     modified and this statement is not removed. Usage by and for commercial
CC
     entities requires a license agreement (See http://www.isb-sib.ch/announce/
CC
     or send an email to license@isb-sib.ch).
CC
CC
     EMBL; A3036829; BAA92340.1; -.
DR
DR
     EMBL; AB036830; BAA92341.1; -.
DR
     EMBL; AB036831; BAA92342.1; -.
     EMBL; BC004362; AAH04362.1; -.
DR
     EMBL; U45973; AAB03214.1; -.
DR
     HSSP; 043001; 119Z.
DR
     MIM; 607875; -.
DR
DR
     GO; GO:0005829; C:cytosol; ISS.
     GO; GO:0043005; C:neuronal cell projection; ISS.
DR
     GO; GO:0016312; F:inositol bisphosphate phosphatase activity; ISS.
DR
     GO; GO:0046030; F:inositol trisphosphate phosphatase activity; ISS.
DR
     GO; GO:0042577; F:lipid phosphatase activity; ISS.
DR
     GC; GO:0030036; P:actin cytoskeleton organization and biogenesis; ISS.
DR
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DR
    InterPro; IPR005135; Exo endo phos.
    InterPro; IPR000300; IPPc.
DR
    Pfam; PF03372; Exo endo phos; 1.
DR
    SMART; SM00128; IPPc; 1.
DR
    Alternative splicing; Endoplasmic reticulum; Hydrolase.
KW
                     318
                              Catalytic (Potential).
FT
    DOMAIN
                16
                     448
                              Required for ruffle localization.
    DOMAIN
               321
FT
    VARSPLIC
                      76
                              Missing (in isoform 2).
FT
                              /FTId=VSP 050612.
FT
                              Y->A,F: No effect on EGF-induced ruffle
FT
    MUTAGEN
               349
                     349
TH
                              localization.
                              D->A: Significant decrease in EGF-induced
FT
    MUTAGEN
               361
                     361
                              ruffle localization.
FT
                              W->A: Significant decrease in EGF-induced
FT
    MUTAGEN
               362
                     362
FT
                              ruffle localization.
                              Y->A,F: No effect on EGF-induced ruffle
FT
    MUTAGEN
               376
                     376
                              localization.
FT
                              T \rightarrow A \text{ (in Ref. 3)}.
FT
    CONFLICT
               120
                     120
                              R \rightarrow S \text{ (in Ref. 2)}.
FT
    CONFLICT
               416
                     416
                     426
FT
    CONFLICT
               426
                              R \rightarrow S \text{ (in Ref. 2)}.
SQ
    SEQUENCE
              448 AA; 51228 MW; 0651BE3C6E2EEB0F CRC64;
 Query Match
                       99.5%;
                              Score 2010; DB 1; Length 448;
 Best Local Similarity
                       99.5%;
                              Pred. No. 1.1e-164;
 Matches 370; Conservative
                             0; Mismatches
                                            2;
                                                Indels
                                                         0; Gaps
          1 MDVLSPLSFIKVSHVRMQGILLLVFAKYQHLPYTQILSTKSTPTGLFGYWGNKGGVNICL 60
Qy
            77 MDVLSPLSFIKVSHVRMQGILLLVFAKYQHLPYIQILSTKSTPTGLFGYWGNKGGVNICL 136
CC
         61 KLYGYYVSIINCHLPPHISNNYQRLEHFDRILEMQNCEGRDIPNILDHDLIIWFGDMNFR 120
QУ
            137 KLYGYYVSIINCHLPPHISNNYORLEHFDRILEMONCEGRDIPNILDHDLIIWFGDMNFR 196
Db
         121 IEDFGLHFVRESIKNRCYGGLWEKDQLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDY 180
Qy
            197 IEDFGLHFVRESIKNRCYGGLWEKDQLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDY 256
Db
         181 DTSEKKRKPAWTDRILWRLKRQPCAGPDTPIPPASHFSLSLRGYSSHMTYGISDHKPVSG 240
QУ
            257 DTSEKKRKPAWTDRILWRLKRQPCAGPDTPIPPASHFSLSLRGYSSHMTYGISDHKPVSG 316
Db
         241 TFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVNDY 300
Qу
            317 TFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVNDY 376
Db
         301 VSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFQIPPGSL 360
Qу
            377 VSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYRNSLRSVVGIRRPFQIPPGSL 436
Db
         361 REDPLGEAQPQI 372
QУ
            Db
         437 REDPLGEAQPQI 448
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RESULT 2 SKIP MOUSE

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SKIP MOUSE
ID
                    STANDARD;
                                    PRT;
                                           468 AA.
     O8C5L6; O09040;
AC
DT
     10-OCT-2003 (Rel. 42, Created)
     29-MAR-2004 (Rel. 43, Last sequence update)
DΤ
     05-JUL-2004 (Rel. 44, Last annotation update)
DT
DΕ
     Skeletal muscle and kidney enriched inositol phosphatase
DΕ
     (EC 3.1.3.56).
GN
     Name=Skip; Synonyms=Pps;
os
     Mus musculus (Mouse).
     Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
     Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
OC
OX
     NCBI TaxID=10090;
RN
     [1]
RP
     SEQUENCE FROM N.A.
     MEDLINE=97325785; PubMed=9182797; DOI=10.1016/S0896-6273(00)80312-8;
RX
     Hamilton B.A., Smith D.J., Mueller K.L., Kerrebrock A.W.,
RA
     Bronson R.T., van Berkel V., Daly M.J., Kruglyak L., Reeve M.P.,
RA
     Nemhauser J.L., Hawkins T.L., Rubin E.M., Lander E.S.;
RA
     "The vibrator mutation causes neurodegeneration via reduced expression
RT
     of PITP alpha: positional complementation cloning and extragenic
RT
RT
     suppression.";
RL
     Neuron 18:711-722(1997).
RN
     [2]
RP
     SEQUENCE FROM N.A.
RC
     STRAIN=ILS, and ISS;
     Ehringer M.A., Thompson J., Conroy O., Xu Y., Yang F., Canniff J.,
RA.
RA
     Beeson M., Gordon L., Bennett B., Johnson T.E., Sikela J.M.;
RL
     Submitted (FEB-2002) to the EMBL/GenBank/DDBJ databases.
RN \cdot
RP
     SEQUENCE FROM N.A.
     STRAIN=C57BL/6J; TISSUE=Medulla oblongata, and Ovary;
RC
RX
     MEDLINE=22354683; PubMed=12466851; DOI=10.1038/nature01266;
Æ.
     Okazaki Y., Furuno M., Kasukawa T., Adachi J., Bono H., Kondo S.,
RА
     Nikaido I., Osato N., Saito R., Suzuki H., Yamanaka I., Kiyosawa H.,
RA
     Yagi K., Tomaru Y., Hasegawa Y., Nogami A., Schonbach C., Gojobori T.,
RA
     Baldarelli R., Hill D.P., Bult C., Hume D.A., Quackenbush J.,
RA
     Schriml L.M., Kanapin A., Matsuda H., Batalov S., Beisel K.W.,
     Blake J.A., Bradt D., Brusic V., Chothia C., Corbani L.E., Cousins S.,
RΑ
     Dalla E., Dragani T.A., Fletcher C.F., Forrest A., Frazer K.S.,
RA.
     Gaasterland T., Gariboldi M., Gissi C., Godzik A., Gough J.,
RΆ
     Grimmond S., Gustincich S., Hirokawa N., Jackson I.J., Jarvis E.D.,
RA
     Kanai A., Kawaji H., Kawasawa Y., Kedzierski R.M., King B.L.,
RA
     Konagaya A., Kurochkin I.V., Lee Y., Lenhard B., Lyons P.A.,
RA
     Maglott D.R., Maltais L., Marchionni L., McKenzie L., Miki H.,
RA
     Nagashima T., Numata K., Okido T., Pavan W.J., Pertea G., Pesole G.,
RA
     Petrovsky N., Pillai R., Pontius J.U., Qi D., Ramachandran S.,
RA
     Ravasi T., Reed J.C., Reed D.J., Reid J., Ring B.Z., Ringwald M.,
RA
     Sandelin A., Schneider C., Semple C.A., Setou M., Shimada K.,
RA
     Sultana R., Takenaka Y., Taylor M.S., Teasdale R.D., Tomita M.,
RA
     Verardo R., Wagner L., Wahlestedt C., Wang Y., Watanabe Y., Wells C.,
RA
     Wilming L.G., Wynshaw-Boris A., Yanagisawa M., Yang I., Yang L.,
RA
     Yuan Z., Zavolan M., Zhu Y., Zimmer A., Carninci P., Hayatsu N.,
RA
     Hirozane-Kishikawa T., Konno H., Nakamura M., Sakazume N., Sato K.,
RΑ
     Shiraki T., Waki K., Kawai J., Aizawa K., Arakawa T., Fukuda S.,
RA
     Hara A., Hashizume W., Imotani K., Ishii Y., Itoh M., Kagawa I.,
RA
     Miyazaki A., Sakai K., Sasaki D., Shibata K., Shinagawa A.,
RA
     Yasunishi A., Yoshino M., Waterston R., Lander E.S., Rogers J.,
RA
```

```
RA
    Birney E., Hayashizaki Y.;
RT
    "Analysis of the mouse transcriptome based on functional annotation of
RT
    60,770 full-length cDNAs.";
    Nature 420:563-573(2002).
RL
    -!- FUNCTION: Inositol 5-phosphatase which acts on inositol 1,4,5-
CC
        trisphosphate, inositol 1,3,4,5-tetrakisphosphate,
CC
CC
        phosphatidylinositol 4,5-bisphosphate and phosphatidylinositol
CC
        3,4,5-triphosphate. Has 6-fold higher affinity for
CC
        phosphatidylinositol 4,5-bisphosphate than for inositol 1,4,5-
CC
        trisphosphate. May negatively regulate assembly of the actin
        cytoskeleton (By similarity).
CC
CC
    -!- CATALYTIC ACTIVITY: D-myo-inositol 1,4,5-trisphosphate + H(2)0 =
CC
        myo-inositol 1,4-bisphosphate + phosphate.
CC
    -!- SUBCELLULAR LOCATION: Endoplasmic reticulum. Following stimulation
CC
        with EGF, translocates to membrane ruffles (By similarity).
CC
    -!- SIMILARITY: Belongs to the inositol-1,4,5-trisphosphate 5-
CC
        phosphatase type II family.
    ______
CC
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CC
CC
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CC.
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    modified and this statement is not removed. Usage by and for commercial
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CC
    entities requires a license agreement (See http://www.isb-sib.ch/announce/
CC
    or send an email to license@isb-sib.ch).
    ______
CC.
DR EMBL; U96724; AAC53265.1; -.
DR
    EMBL; U96726; AAC60757.1; -.
    EMBL; AF483522; AAL90796.1; -.
DR
    EMBL; AF483523; AAL90797.1; -.
DR
    EMBL; AK054436; BAC35778.1; -.
DR
DR
    EMBL; AK078104; BAC37126.1; -.
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    MGD; MGI:1194899; Pps.
DR
    GO; GO:0005829; C:cytosol; ISS.
DR
    GO; GO:0043005; C:neuronal cell projection; ISS.
DR
    GO; GO:0016312; F:inositol bisphosphate phosphatase activity; ISS.
DR
    CO; GO:0046030; F:inositol trisphosphate phosphatase activity; ISS.
DR
DR.
    GO; GO:0042577; F:lipid phosphatase activity; ISS.
DR
    GO; GO:0030036; P:actin cytoskeleton organization and biogenesis; ISS.
DR
    InterPro; IPR005135; Exo endo phos.
DR
    InterPro; IPR000300; IPPc.
DR
    Pfam; PF03372; Exo endo phos; 1.
    SMART; SM00128; IPPc; 1.
DR
    Endoplasmic reticulum; Hydrolase.
KN
                               Catalytic (Potential).
FT
    DOMAIN
                34
                      337
FT. DOMAIN
               340
                      468
                               Required for ruffle localization.
                              D -> E (in Ref. 3; BAC37126).
FT
               329
                     329
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SO SEQUENCE
               468 AA; 54158 MW; F2E1CA370B97A8A1 CRC64;
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 Matches 283; Conservative 39; Mismatches
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Qу
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          95 MDMLSPLNFVKISQVRMQGLLLLVFAKYQHLPYIQIISTKSTPTGLYGYWGNKGGVNVCL 154
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61 KLYGYYVSIINCHLPPHISNNYORLEHFDRILEMONCEGRDIPNILDHDLIIWFGDMNFR 120
Qу
            155 KLYGYYVSIINCHLPPHMYNNDQRLEHFDRILESLTFEGYDVPNILDHDLILWFGDMNFR 214
Db
        121 IEDFGLHFVRESIKNRCYGGLWEKDQLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDY 180
QУ
            215 IEDFGLLFVQESITRKYYKELWEKDQLFIAKKNDQLLREFQEGPLLFPPTYKFDRHSNNY 274
Db
        181 DTSEKKRKPAWTDRILWRLKROPC-AGPDTPIPPASHFSLSLRGYSSHMTYGISDHKPVS 239
Qу
            275 DTSEKKRKPAWTDRILWRLKROPSOASPLASSVPTSYFLLTLKNYVSHMAYSISDHKPVT 334
Db
        240 GTFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVND 299
Qу
            335 GTFDLELNPLMSVPLITMMPEHLWTMENDMLISYTSTPEFLSSSWDWIGLYKVGMRHIND 394
Db
QУ
        300 YVSYAWVGDSKVSCSDNLNOVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFQIPPGS 359
            395 YVAYVWVGDNQVSYGNNPNQVYINISAIPDTEDQFLLCYYSNNLHSVVGISQPFKIPIRS 454
Db
Q٧٠
        360 -LREDPLGEAQPQI 372
             455 ELREDTLYEPEPOI 468
Db
RESULT 3
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ID
    Q6CM99
               PRELIMINARY:
                               PRT:
AC
    26GN22;
    05-JUL-2004 (TrEMBLrel. 27, Created)
DT
    J5-JUL-2004 (TrEMBLrel. 27, Last sequence update)
DT
D''
    05-JUL-2004 (TrEMBLrel. 27, Last annotation update)
    LCC443677 protein (Fragment).
DΞ
GN
    Name=LOC443677;
    Xenopus laevis (African clawed frog).
OS
OC
    Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
    Amphibia; Batrachia; Anura; Mesobatrachia; Pipoidea; Pipidae;
CC
OC
    Kenopodinae; Kenopus.
XC
   NCBI TaxID=8355;
RN
    [1]
RP
    SEQUENCE FROM N.A.
    TISSUE=Embryo;
RC
    MEDILINE=22388257; PubMed=12477932; DOI=10.1073/pnas.242603899;
RX
    Strausberg R.L., Feingold E.A., Grouse L.H., Derge J.G.,
RZ
    Klausner R.D., Collins F.S., Wagner L., Shenmen C.M., Schuler G.D.,
RA
    Altschul S.F., Zeeberg B., Buetow K.H., Schaefer C.F., Bhat N.K.,
RS
    Hopkins R.F., Jordan H., Moore T., Max S.I., Wang J., Hsieh F.,
RA
    Diatchenko L., Marusina K., Farmer A.A., Rubin G.M., Hong L.,
RA
    Stapleton M., Soares M.B., Bonaldo M.F., Casavant T.L., Scheetz T.E.,
RA
    Brownstein M.J., Usdin T.B., Toshiyuki S., Carninci P., Prange C.,
RA
    Raha S.S., Loquellano N.A., Peters G.J., Abramson R.D., Mullahy S.J.,
RA
    Bosak S.A., McEwan P.J., McKernan K.J., Malek J.A., Gunaratne P.H.,
RA
    Richards S., Worley K.C., Hale S., Garcia A.M., Gay L.J., Hulyk S.W.,
RA
    Villalon D.K., Muzny D.M., Sodergren E.J., Lu X., Gibbs R.A.,
RA
    Fahey J., Helton E., Ketteman M., Madan A., Rodriques S., Sanchez A.,
RA
    Whiting M., Madan A., Young A.C., Shevchenko Y., Bouffard G.G.,
RA
```

```
Blakesley R.W., Touchman J.W., Green E.D., Dickson M.C.,
RA
    Rodriquez A.C., Grimwood J., Schmutz J., Myers R.M., Butterfield Y.S.,
RA
    Krzywinski M.I., Skalska U., Smailus D.E., Schnerch A., Schein J.E.,
RA
    Jones S.J., Marra M.A.;
RA
    "Generation and initial analysis of more than 15,000 full-length human
RT
    and mouse cDNA sequences.";
RT
    Proc. Natl. Acad. Sci. U.S.A. 99:16899-16903(2002).
RL
RN
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RP
RC
    TISSUE=Embryo;
    MEDLINE=22341132; PubMed=12454917; DOI=10.1002/dvdy.10174;
RX
    Klein S.L., Strausberg R.L., Wagner L., Pontius J., Clifton S.W.,
RA
RA
    Richardson P.;
RT
    "Genetic and genomic tools for Xenopus research: The NIH Xenopus
RT
    initiative.";
    Dev. Dyn. 225:384-391(2002).
RL
RN
    [3]
RP
    SEQUENCE FROM N.A.
RC
    TISSUE=Embryo;
    Klein S., Gerhard D.S.;
RA
RL
    Submitted (JUN-2004) to the EMBL/GenBank/DDBJ databases.
DR
    EMBL; BC073616; AAH73616.1; -.
    GO; GO:0004437; F:inositol or phosphatidylinositol phosphatas. . .; IEA.
DR
    InterPro; IPR000345; CytC_heme BS.
DR
    InterPro; IPR005135; Exo endo phos.
DR
DR
    InterPro; IPR000300; IPPc.
    Pfam; PF03372; Exo endo phos; 1.
DR
DR
    SMART; SM00128; IPPc; 1.
    PROSITE; PS00190; CYTOCHROME C; UNKNOWN 1.
DR
FT
    NON TER
                 1
                       1
SO
    SEQUENCE
              496 AA; 56894 MW; 832B0717978C7D63 CRC64;
  Query Match
                       55.2%; Score 1115; DB 2; Length 496;
                       53.1%; Pred. No. 1.9e-87;
  Best Local Similarity
  Matches 203; Conservative 65; Mismatches 100; Indels
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Qу
             112 MDILAPHGYVKLSSIRLOGLLLLTFVKHOHIPFVQDIRTNYIRTGLFGYWGNKGGVTVRM 171
Db
          61 KLYGYYVSIINCHLPPHISNNYORLEHFDRILEMONCEGRDIPNILDHDLIIWFGDMNFR 120
QУ
              172 SVYGHMICFMNCHLPAHMENTNORLDDFERMLDTOOFDDDNTGNILDHDLVFCFGDLNFR 231
Db
         121 IEDFGLHFVRESIKNRCYGGLWEKDQLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDY 180
Qу
             232 ITDFGIHFIREAINSNKYNLLWEKDQLNIAKKKEQFLQGFLEGPLKFKPTYKFDQNSDVY 291
Db
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Qу
             -:||--|:|||:|||:|||
         2)2 DTSSKKRKPAWTDRILWKLK--DVSGSDPGETTGSDFEEFLKVSLDKYTSHMSYGISDHK 349
Db
         237 PVSGTFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRD 296
Qу
             350 PVTGTFTLQLKPLILTPSVTLKPEGEWNADHDSLISYSVAKDFPSSTWDWIGLYRVGFRH 409
Db
         297 VNDYVSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFQIP 356
Qу
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410 TNDYVTYAWVKDDEISFSEDCNQVYMNADDIPLDGGEFLLCYHCHNLQCLAGISQPFQIR 469
Db
          357 PG-----SLREDPLGEAQP 370
Qу
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          470 PGRNLGLKEEENRESPFKTESP 491
Db
RESULT 4
P5PA MOUSE
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                    STANDARD;
                                   PRT;
                                         1003 AA.
AC
     P59644;
DT
     10-OCT-2003 (Rel. 42, Created)
DT
     10-OCT-2003 (Rel. 42, Last sequence update)
DT
     05-JUL-2004 (Rel. 44, Last annotation update)
     Fhosphatidylinositol 4,5-bisphosphate 5-phosphatase A (EC 3.1.3.56).
DE
GN
     Name=Pib5pa;
os
     Mus musculus (Mouse).
     Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
OC
     Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
OX
     NCBI TaxID=10090;
RN
     [1]
RP
     SEQUENCE FROM N.A.
     STRAIN=C57BL/6J; TISSUE=Diencephalon;
RC
     MEDLINE=22354683; PubMed=12466851; DOI=10.1038/nature01266;
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     Okazaki Y., Furuno M., Kasukawa T., Adachi J., Bono H., Kondo S.,
RA
     Nikaido I., Osato N., Saito R., Suzuki H., Yamanaka I., Kiyosawa H.,
\mathbb{R}^{N}
     Yaqi K., Tomaru Y., Haseqawa Y., Noqami A., Schonbach C., Gojobori T.,
RA
     Daldarelli R., Hill D.P., Bult C., Hume D.A., Quackenbush J.,
RΑ
     Schriml L.M., Kanapin A., Matsuda H., Batalov S., Beisel K.W.,
RA
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     Dalla E., Dragani T.A., Fletcher C.F., Forrest A., Frazer K.S.,
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     Gaasterland T., Gariboldi M., Gissi C., Godzik A., Gough J.,
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     Grimmond S., Gustincich S., Hirokawa N., Jackson I.J., Jarvis E.D.,
RA
     Kanai A., Kawaji H., Kawasawa Y., Kedzierski R.M., King B.L.,
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     Konagaya A., Kurochkin I.V., Lee Y., Lenhard B., Lyons P.A.,
RA
     Maglott D.R., Maltais L., Marchionni L., McKenzie L., Miki H.,
RA
     Hajashima T., Numata K., Okido T., Pavan W.J., Pertea G., Pesole G.,
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     Petrovsky N., Pillai R., Pontius J.U., Qi D., Ramachandran S.,
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     Ravasi T., Reed J.C., Reed D.J., Reid J., Ring B.Z., Ringwald M.,
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     Sandelin A., Schneider C., Semple C.A., Setou M., Shimada K.,
RA
     Sultana R., Takenaka Y., Taylor M.S., Teasdale R.D., Tomita M.,
RA
     Verardo R., Wagner L., Wahlestedt C., Wang Y., Watanabe Y., Wells C.,
RA
     Wilming L.G., Wynshaw-Boris A., Yanagisawa M., Yang I., Yang L.,
RA
     Yuan Z., Zavolan M., Zhu Y., Zimmer A., Carninci P., Hayatsu N.,
RA
     Hirozane-Kishikawa T., Konno H., Nakamura M., Sakazume N., Sato K.;
RA
     Shiraki T., Waki K., Kawai J., Aizawa K., Arakawa T., Fukuda S.,
RA
     Hara A., Hashizume W., Imotani K., Ishii Y., Itoh M., Kagawa I.,
RA.
     Miyazaki A., Sakai K., Sasaki D., Shibata K., Shinagawa A.,
ŖА
     Yasunishi A., Yoshino M., Waterston R., Lander E.S., Rogers J.,
RA
     Dirney E., Hayashizaki Y.;
\mathbf{F}\mathbf{A}
     "Analysis of the mouse transcriptome based on functional annotation of
RT'
     60,770 full-length cDNAs.";
RT
     Nature 420:563-573(2002).
RL
RN
     SUBCELLULAR LOCATION, AND IDENTIFICATION OF DOMAIN REQUIRED FOR
RP
     MEMBRANE RUFFLE LOCALIZATION.
```

RP

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MEDLINE=22538472; PubMed=12536145; DOI=10.1074/jbc.M209991200;
RX
    Gurung R., Tan A., Ooms L.M., McGrath M.J., Huysmans R.D.,
RA
    Munday A.D., Prescott M., Whisstock J.C., Mitchell C.A.;
RA
     "Identification of a novel domain in two mammalian inositol-
RT
    polyphosphate 5-phosphatases that mediates membrane ruffle
RT
    localization. The inositol 5-phosphatase SKIP localizes to the
RT
    endoplasmic reticulum and translocates to membrane ruffles following
RT
    epidermal growth factor stimulation.";
RT
    J. Biol. Chem. 278:11376-11385(2003).
RL
    -!- FUNCTION: Inositol 5-phosphatase, which converts inositol 1,4,5-
CC
CC
        trisphosphate to inositol 1,4-bisphosphate. Also converts
CC
        phosphatidylinositol 4,5-bisphosphate to phosphatidylinositol 4-
CC
        phosphate and inositol 1,3,4,5-tetrakisphosphate to inositol
CC
        1,3,4-trisphosphate in vitro. May be involved in modulation of the
CC
        function of inositol and phosphatidylinositol polyphosphate-
CC
        binding proteins that are present at membranes ruffles (By
CC
        similarity).
CC
     -!- CATALYTIC ACTIVITY: D-myo-inositol 1,4,5-trisphosphate + H(2)O =
CC
        myo-inositol 1,4-bisphosphate + phosphate.
CC
    -!- CATALYTIC ACTIVITY: 1D-myo-inositol 1,3,4,5-tetrakisphosphate +
CC
        H(2)O = 1D-myo-inositol 1,3,4-trisphosphate + phosphate.
CC
    -!- SUBCELLULAR LOCATION: Cytoplasmic; predominantly localized to
CC
        membrane ruffles.
CC
    -!- DOMAIN: The 5 Arg-Ser-Xaa-Ser-Xaa-Xaa (RSXSXX) motifs may
CC
        constitute binding sites for the 14-3-3 protein.
CC
    -!- PTM: Phosphorylated at Ser/Thr residues (By similarity).
    -!- SIMILARITY: Belongs to the inositol-1,4,5-trisphosphate 5-
CC
C_{1}
     phosphatase type II family.
CC
    ______
CC
    This SWISS-PROT entry is copyright. It is produced through a collaboration
CC
    between the Swiss Institute of Bioinformatics and the EMBL outstation -
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    the European Bioinformatics Institute. There are no restrictions on its
CC
    use by non-profit institutions as long as its content is in no way
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    modified and this statement is not removed. Usage by and for commercial
CC
    entities requires a license agreement (See http://www.isb-sib.ch/announce/.
CC
    or send an email to license@isb-sib.ch).
CC
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    EMBL; AK034272; BAC28654.1; -.
ЭR
DR
    HSSP; 043001; 119Z.
DR
    MGD; MGI:2158663; Pib5pa.
DR
    InterPro; IPR005135; Exo endo phos.
DR
    InterPro; IPRC00300; IPPc.
DR
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    JMART; SM00128; IPPC; 1.
DR
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ΚW
FT
                                Catalytic (Potential).
    MIAMOD
                422
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FΤ
    DOMAIII
                726
                       837
                                Required for ruffle localization.
FT
    DOMAIN
                50
                       374
                                Pro-rich.
FT
    · DOMAIN
                837
                       934
                                Ser-rich.
    SITE
                                RSXSXX motif 1.
FT
                 6
                       11
                                RSXSXX motif 2.
                351
                       356
FT
    SITE
FT
    SITE
                871
                       876
                                RSXSXX motif 3.
F''
    SITE
                882
                       887
                                RSXSXX motif 4.
    SITE
                908
                                RSXSXX motif 5.
FT
                      913
FT
                                SH3-binding (Potential).
    SITE
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               1003 AA; 107602 MW; AFF4FD929CFCB6BA CRC64;
SQ
    SEQUENCE
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Db
          61 KLYGYYVSIINCHLPPHISNNYORLEHFDRILEMONCEGRDIPNILDHDLIIWFGDMNFR 120
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Db
         121 IEDFGLHFVRESIKNRCYGGLWEKDQLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDY 180
Qy
             || : ||||: :| :
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         181 DTSEKKRKPAWTDRILWRLKRQPCAGPDTPIPPASH-FSLSLRGYSSHMTYGISDHKPVS 239
Qу
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Db
         240 GTFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVND 299
Qy
                        723 AQFILQFATRDDVPLVRLEVADEWARPEQAVVRYRVETVFARSSWDWIGLYRVGFRHCKD 782
ďQ
         300 YVSYAWVGDSKVSCSDNLNOVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRFFOI 355
Ç7
             : |
                          |: ||
                                   ::| -
                                          : | : | | | | :
                                                      :: | :: | | | |
         733 YVAYVWAKHEEV--DGNIYQVTFSEESLPKGHGDFILGYYSHHHSILIGVTEPFQI 836
D'S
RESULT 5
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ID
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                  STANDARD;
                                FRT;
                                     1001 AA.
AC
    Q9JMC1;
DT
    10-CCT-2003 (Rel. 42, Created)
    10-OCT-2003 (Rel. 42, Last sequence update)
DT
DT
    05-JUL-2004 (Rel. 44, Last annotation update)
DE
    Phosphatidylinositol 4,5-bisphosphate 5-phosphatase A (EC 3.1.3.56)
DE
    (Proline-rich inositol polyphosphate 5-phosphatase).
GN
    Name=Pib5pa; Synonyms=Pipp;
os
    Rattus norvegicus (Rat).
    Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC.
OC
    Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Rattus.
OX
    NCBI TaxID=10116;
RN
    SEQUENCE FROM N.A., SUBCELLULAR LOCATION, TISSUE SPECIFICITY, AND
RP
R۵
    ENZYMATIC ACTIVITY.
RC
    TISSUE=Brain:
RX
    MEDLINE=20062911; PubMed=10593988; DOI=10.1074/jbc.274.51.36790;
RA
    Mochizuki Y., Takenawa T.;
RT
    "Novel inositol polyphosphate 5-phosphatase localizes at membrane
RT
    ruffles.";
    J. Biol. Chem. 274:36790-36795(1999).
RI_{i}
    -!- FUNCTION: Inositol 5-phosphatase, which converts inositol 1,4,5-
CC
CC
        trisphosphate to inositol 1,4-bisphosphate. Also converts
CC
        phosphatidylinositol 4,5-bisphosphate to phosphatidylinositol 4-
CC
        phosphate and inositol 1,3,4,5-tetrakisphosphate to inositol
CC
        1,3,4-trisphosphate in vitro. May be involved in modulation of the
```

```
CC
        binding proteins that are present at membranes ruffles.
    -!- CATALYTIC ACTIVITY: D-myo-inositol 1,4,5-trisphosphate + H(2)0 =
CC
        myo-inositol 1,4-bisphosphate + phosphate.
CC
    -!- CATALYTIC ACTIVITY: 1D-myo-inositol 1,3,4,5-tetrakisphosphate +
CC
        H(2)O = 1D-myo-inositol 1,3,4-trisphosphate + phosphate.
CC
    -!- SUBCELLULAR LOCATION: Cytoplasmic; predominantly localized to
CC
CC
        membrane ruffles.
    -!- TISSUE SPECIFICITY: Expressed in heart, brain, kidney, stomach,
CC
        small intestine and lung. Not expressed in spleen, thymus,
CC
        skeletal muscle, testis and skin.
CC
    -!- DOMAIN: The 5 Arg-Ser-Xaa-Ser-Xaa-Xaa (RSXSXX) motifs may
CC
        constitute binding sites for the 14-3-3 protein.
CC
    -!- PTM: Phosphorylated on Ser/Thr residues.
CC
    -!- SIMILARITY: Belongs to the inositol-1,4,5-trisphosphate 5-
CC
CC
        phosphatase type II family.
    ______
CC
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CC
    between the Swiss Institute of Bioinformatics and the EMBL outstation -
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    the European Bioinformatics Institute. There are no restrictions on its
CC
    use by non-profit institutions as long as its content is in no way
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    modified and this statement is not removed. Usage by and for commercial
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    entities requires a license agreement (See http://www.isb-sib.ch/announce/
CC
    or send an email to license@isb-sib.ch).
CC
    CC
DR
    EMBL; AB032551; BAA90553.1; -.
DR '
    HSSP, 043001; 119Z.
DR
    InterPro; IPR005135; Exo endo phos.
    InterPro; IPR000300; IPPc.
DR
    Pfam; PF03372; Exo endo phos; 1.
DR
    SMART; SM00128; IPPC; 1.
DR
KW
    Hydrolase; Phosphorylation; Repeat; SH3-binding.
FT
    MILAMOC
               420
                      723
                               Catalytic (Potential).
FT
    DOMAIN
               724
                      835
                               Required for ruffle localization (By
FT
                               similarity).
               50
                      387
                               Pro-rich.
FT
    NIAMOG
                      932
                               Ser-rich.
FT
    DOMAIN
               835
                               RSXSXX motif 1.
                      11
FT
    SITE
                6
                               RSXSXX motif 2.
FT
    SITE
               351
                      356
    SITE
FT
               869
                      874
                               RSXSXX motif 3.
FT
               880
                    885
                               RSXSXX motif 4.
    SITE
FT
    SITE
               906
                     911
                               RSXSXX motif 5.
                               SH3-binding (Potential).
FT
               346
                     351
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    SEQUENCE
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                       43.2%; Score 872.5; DB 1; Length 1001;
 Query Match
 Best Local Similarity 47.5%; Pred. No. 4e-66;
  Matches 169; Conservative 56; Mismatches 126; Indels
                                                           5; Gaps
                                                                      4;
           1 MDVLSPLSFIKVSHVRMQGILLLVFAKYQHLPYIQILSTKSTPTGLFGYWGNKGGVNICL 60
Qу
             483 MDALGPFNFVLVSTVRMQGVILLLFAKYYHLPFLRDVQTDCTRTGLGGYWGNKGGVSVRL 542
Db
          61 KLYGYYVSIINCHLPPHISNNYQRLEHFDRILEMQNCEGRDIPNILDHDLIIWFGDMNFR 120
Qу
                                                    || ::| || :| :|
               :|: : :|||| |:
         543 AAFGHMLCFLNCHLPAHMDKAEQRKDNFQTILSLQQFQGPGAHGILDHDLVFWFGDLNFR 602
Db
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function of inositol and phosphatidylinositol polyphosphate-

CC

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121 IEDFGLHFVRESIKNRCYGGLWEKDQLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDY 180
Qу
             || : |||| : | :
                               - |:|: |||i | | ||:||| :| |
         603 IESYDLHFVKFAIDSNQLHQLWEKDQLNMAKNTWPILKGFQEGPLNFAPTFKFDVGTNKY 662
Dh
         181 DTSEKKRKPAWTDRILWRLKRQPCAGPDTPIPPASH-FSLSLRGYSSHMTYGISDHKPVS 239
Qу
             663 DTSAKKKPAWTDRILWKVK-APSGGP-SPSGRESHRLQVTQHSYRSHMEYTVSDHKPVA 720
Db
         240 GTFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVND 299
Qу
              721 ARFLLOFAFRDDVPLVRLEVADEWARPEOAVVRYRVETVFARSSWDWIGLYRVGFRHCKD 780
Db
         300 YVSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFQI 355
Qу
             1|:| | :| |: || ::| :|:| |||
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Db
RESULT 6
O6ZTH6
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ID
    Q6ZTH6
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AC
    Q6ZTH6;
DT
    05-JUL-2004 (TrEMBLrel. 27, Created)
    05-JUL-2004 (TrEMBLrel. 27, Last sequence update)
DT
    05-JUL-2004 (TrEMBLrel. 27, Last annotation update)
ÐΤ
DΕ
    Hypothetical protein FLJ44647.
ാട
    Homo saviens (Human).
    Cukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
    Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
OC
OX
    MCBI TaxID=9606;
327
    [1]
RΡ
    SEQUENCE FROM N.A.
RC
    TISSUE=Cerebellum;
RA
    Tanigami A., Fujiwara T., Shibahara T., Goto Y., Hirao M., Shimizu F.,
RA
    Wakebe H., Ono T., Hishigaki H., Watanabe T., Ozaki K., Sugiyama T.,
    Trie R., Otsuki T., Sato H., Wakamatsu A., Ishii S., Yamamoto J.,
SV
    Isono Y., Kawai-Hio Y., Saito K., Nishikawa T., Kimura K.,
RA
    Yamashita H., Matsuo K., Nakamura Y., Sekine M., Kikuchi H., Kanda K.,
RA
    Wagatsuma M., Murakawa K., Kanehori K., Takahashi-Fujii A., Oshima A.,
RA
    Suqiyama A., Kawakami B., Suzuki Y., Sugano S., Nagahari K.,
A.7
RA
    Masuho Y., Nagai K., Isogai T.;
    Submitted (JUL-2003) to the EMBL/GenBank/DDBJ databases.
RL
\mathbb{DR}
    EMBL; AK126610; BAC86611.1; -.
    GO; GO:0004437; F:inositol or phosphatidylinositol phosphatas. . .; IEA.
DR
    InterPro; IPR005135; Exo endo phos.
DR
    InterPro; IPR000300; IPPc.
DR
    Pfam; PF03372; Exo endo phos; 1.
\mathtt{DR}
    SMART; SM00128; IPPc; 1.
DR
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  Query Match
                        47.8%; Pred. No. 3.3e-66;
 Best Local Similarity
 Matches 170; Conservative 54; Mismatches 127; Indels
                                                            5; Gaps
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Qу
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Db
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                                   || ::| || :| :|
                                                        180 AAFGHMLCFLNCHLPAHMDKAEQRKDNFQTILSLQQFQGPGAQGILDHDLVFWFGDLNFR 239
Db
          121 IEDFGLHFVRESIKNRCYGGLWEKDQLSIAKKHDPLLREFQEGRLLFPPTYKFDRNSNDY 180
Qу
                                 111111::11
                                              || : ||||: :| :
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Db
         240 GTFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVND 299
Qy
                          Db
          358 AOFLLOFAFRDDMPLVRLEVADEWVRPEOAVVRYRMETVFARSSWDWIGLYRVGFRHCKD 417
         300 YVSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFQI 355
Qу
                     ::| :|:| |||::
                                                         ::||: ||||
Db
         418 YVAYVWAKHEDV--DGNTYQVTFSEESLPKGHGDFILGYYSHNHSILIGITEPFQI 471
RESULT 7
MAMUH AGEG
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     10-OCT-2003 (Rel. 42, Created)
ra
     10-CCT-2003 (Rel. 42, Last sequence update)
DT.
     05-JUL-2004 (Rel. 44, Last annotation update)
     Phosphatidylinositol 4,5-bisphosphate 5-phosphatase A (EC 3.1.3.56):
DΕ
GN
     Name=PIB5PA; Synonyms=PIPP;
OS
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OC
     Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
     Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
\Sigma
     NUBI_TaxID=9606;
RN
     SEQUENCE FROM N.A. (ISOFORM 2).
RP
RC
     TISSUE=Heart;
     PubMed=14702039; DOI=10.1038/ng1285;
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RΑ
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Fujimori Y., Komiyama M., Tashiro H., Taniqami A., Fujiwara T.,
RA
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     Okumura K., Nagase T., Nomura N., Kikuchi H., Masuho Y., Yamashita R.,
RA
     Nakai K., Yada T., Nakamura Y., Ohara O., Isogai T., Sugano S.;
RA
     "Complete sequencing and characterization of 21,243 full-length human
RT
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     cDNAs.";
     Nat. Genet. 36:40-45(2004).
RL
RN
RP
     SEQUENCE FROM N.A. (ISOFORM 1).
RX
     MEDLINE=20057165; PubMed=10591208; DOI=10.1038/990031;
     Dunham I., Hunt A.R., Collins J.E., Bruskiewich R., Beare D.M.,
RA
     Clamp M., Smink L.J., Ainscough R., Almeida J.P., Babbage A.K.,
RĄ
RA
     Bagguley C., Bailey J., Barlow K.F., Bates K.N., Beasley O.P.,
     Bird C.P., Blakey S.E., Bridgeman A.M., Buck D., Burgess J.,
RA
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RA
     Clegg S.M., Cobley V.E., Cole C.G., Collier R.E., Connor R.,
RA
RΑ
     Conroy D., Corby N.R., Coville G.J., Cox A.V., Davis J., Dawson E.,
     Dhami P.D., Dockree C., Dodsworth S.J., Durbin R.M., Ellington A.G.,
RA
     Lvans K.L., Fey J.M., Fleming K., French L., Garner A.A.,
RA
     Cilbert J.G.R., Goward M.E., Grafham D.V., Griffiths M.N.D., Hall C.,
RA
     Hall R.E., Hall-Tamlyn G., Heathcott R.W., Ho S., Holmes S.,
RA
     Hunt S.E., Jones M.C., Kershaw J., Kimberley A.M., King A.,
RA
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RA
     Martyn I.D., Mashreghi-Mohammadi M., Matthews L.H., Mccann O.T.,
RA
     Mcclay J., Mclaren S., McMurray A.A., Milne S.A., Mortimore B.J.,
RA
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RA
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RΑ
     Scott C.E., Sehra H.K., Skuce C.D., Smalley S., Smith M.L.,
PA
     Scderlund C., Spragon L., Steward C.A., Sulston J.E., Swann R.M.,
RA
     Vaudin M., Wall M., Wallis J.M., Whiteley M.N., Willey D.L.,
RA
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RA
     Wright C.L., Hubbard T., Bentley D.R., Beck S., Rogers J., Shimizu N.,
RA
     Minoshima S., Kawasaki K., Sasaki T., Asakawa S., Kudoh J.,
R\Lambda
     Shintani A., Shibuya K., Yoshizaki Y., Aoki N., Mitsuyama S.,
RΑ
     Roe B.A., Chen F., Chu L., Crabtree J., Deschamps S., Do A., Do T.,
RA
     Dorman A., Fang F., Fu Y., Hu P., Hua A., Kenton S., Lai H., Lao H.I.,
RA
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ŖА
     Phan S., Qi S., Qian Y., Ray L., Ren Q., Shaull S., Sloan D., Song L.,
RA
     Wang Q., Wang Y., Wang Z., White J., Willingham D., Wu H., Yao Z.,
RA
     Zhan M., Zhang G., Chissoe S., Murray J., Miller N., Minx P.,
RA
RΑ
     Fulton R., Johnson D., Bemis G., Bentley D., Bradshaw H., Bourne S.,
     Cordes M., Du Z., Fulton L., Goela D., Graves T., Hawkins J.,
RA
     Hinds K., Kemp K., Latreille P., Layman D., Ozersky P., Rohlfing T.,
RA
RA
     Schoet P., Walker C., Wamsley A., Wohldmann P., Pepin K., Nelson J.,
     Korf I., Bedell J.A., Hillier L.W., Mardis E., Waterston R.,
RA
RA
     Wilson R., Emanuel B.S., Shaikh T., Kurahashi H., Saitta S.,
     Budarf M.L., McDermid H.E., Johnson A., Wong A.C.C., Morrow B.E.,
RΆ
     Edelmann L., Kim U.J., Shizuya H., Simon M.I., Dumanski J.P.,
RA
RΑ
     Peyrard M., Kedra D., Seroussi E., Fransson I., Tapia I., Bruder C.E.,
RA
     O'Brien K.F., Wilkinson P., Bodenteich A., Hartman K., Hu X.,
RA
     Khan A.S., Lane L., Tilahun Y., Wright H.;
RT
     "The DNA sequence of human chromosome 22.";
```

```
Nature 402:489-495(1999).
RL
RN
     SEQUENCE OF 648-1044 FROM N.A.
RP
     TISSUE=Brain;
RC
RA
     Nussbaum R.L.;
RL
     Submitted (JAN-1996) to the EMBL/GenBank/DDBJ databases.
     -!- FUNCTION: Inositol 5-phosphatase, which converts inositol 1,4,5-
CC
CC
         trisphosphate to inositol 1,4-bisphosphate. Also converts
CC
        phosphatidylinositol 4,5-bisphosphate to phosphatidylinositol 4-
        phosphate and inositol 1,3,4,5-tetrakisphosphate to inositol
CC
         1,3,4-trisphosphate in vitro. May be involved in modulation of the
CC
        function of inositol and phosphatidylinositol polyphosphate-
CC
        binding proteins that are present at membranes ruffles (By
CC
CC
         similarity).
CC
     -!- CATALYTIC ACTIVITY: D-myo-inositol 1,4,5-trisphosphate + H(2)C =
CC
        myo-inositol 1,4-bisphosphate + phosphate.
     -!- CATALYTIC ACTIVITY: 1D-myo-inositol 1,3,4,5-tetrakisphosphate +
CC
CC
        H(2)O = 1D-myo-inositol 1,3,4-trisphosphate + phosphate.
     -!- SUBCELLULAR LOCATION: Cytoplasmic; predominantly localized to
CC
        membrane ruffles (By similarity).
CC
     -!- ALTERNATIVE PRODUCTS:
CC
CC
        Event=Alternative splicing; Named isoforms=2;
CC
CC
          IsoId=Q15735-1; Sequence=Displayed;
CC.
        Name=2:
CC
          IsoId=Q15735-2; Sequence=VSP 007296;
CC .
     -!- DOMAIN: The 5 Arg-Ser-Xaa-Ser-Xaa-Xaa (RSXSXX) motifs may
CC
        constitute binding sites for the 14-3-3 protein.
     -!- PTM: Phosphorylated at Ser/Thr residues (By similarity).
CC
     -!- SIMILARITY: Belongs to the inositol-1,4,5-trisphosphate 5-
CC
CC
        phosphatase type II family.
     -!- CAUTION: Ref.2 sequence differs from that shown due to arroneous
CC
CC
        gene model prediction.
     CC
CC
     This SWISS-PROT entry is copyright. It is produced through a collaboration
CC
     between the Swiss Institute of Bioinformatics and the EMBL outstation -
CC
     the European Bioinformatics Institute. There are no restrictions on its
     use by non-profit institutions as long as its content is in no way
CC
     modified and this statement is not removed. Usage by and for commercial
CC
     entities requires a license agreement (See http://www.isb-sib.ch/announce/
CC
CC
     or send an email to license@isb-sib.ch).
     _____
CC
DR
     EMBL; AK095944; BAC04657.1; -.
     EMBL; AC005005; AAD15618.1; ALT SEQ.
DR
DR
     EMBL; U45975; AAB03216.1; -.
DR
     HSSP; O43001; 119Z.
DR
     Genew; HGNC:8956; PIB5PA.
DR
     MIM; 606481; -.
     InterPro; IPR005135; Exo_endo_phos.
DR
     InterPro; IPR000300; IPPc.
DR
     Pfam; PF03372; Exo endo phos; 1.
DR
DR
     SMART; SM00128; IPPc; 1.
KU
     Alternative splicing; Hydrolase; Phosphorylation; Repeat; SH3-binding.
FT
                425
                       728
                                Catalytic (Potential).
     DOMAIN
                729
FT
     NIAMOG
                       840
                                Required for ruffle localization (By
FT
                                 similarity).
                       371
                                Pro-rich.
FT
     DOMAIN
                122
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DOMAIN
                840
                       937
                                 Ser-rich.
FT
                                 RSXSXX motif 1.
                102
                       107
FT
     SITE
                                 RSXSXX motif 2.
                350
                       355
     SITE
FT
                                RSXSXX motif 3.
 FT
     SITE
                874
                       879
                                 RSXSXX motif 4.
 FT
     SITE
                885
                       890
     SITE
                                RSXSXX motif 5.
                911
                       916
FT
                                 SH3-binding (Potential).
FT
     SITE
                345
                       350
                                Missing (in isoform 2).
FT
     VARSPLIC
                  1
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                                 /FTId=VSP 007296.
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                                 E \rightarrow D (in Ref. 1).
 FT
     CONFLICT
                596
                       596
                                 SYD -> ARG (in Ref. 3).
 FT
     CONFLICT
                610
                       612
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  Query Match
  Best Local Similarity
                         47.5%; Pred. No. 2e-65;
                                                              5; Gaps
  Matches 169; Conservative 55; Mismatches 127; Indels
            1 MDVLSPLSFIKVSHVRMQGILLLVFAKYQHLPYIQILSTKSTPTGLFGYWGNKGGVNICL 60
Qу
              488 MDALGPFNFVLVSSVRMQGVILLLFAKYYHLPFLRDVQTDCTRTGLGGYWGNKGGVSVRL 547
Db
           61 KLYGYYVSIINCHLPPHISNNYQRLEHFDRILEMQNCEGRDIPNILDHDLIIWFGDMNFR 120
 Qу
                :|: : :||||| |: | || ::| || :| :|
                                                       1|||:|: ||||:||
          548 AAFGHMLCFLNCHLPAHMDKAEQRKDNFQTILSLQQFQGPGAQGILDHELVFWFGDLNFR 607
 Db
          121 IEDFGLHFVRESIKNRCYGGLWEKDOLSIAKKHDPLLREFOEGRLLFPPTYKFDRNSNDY 180
 QУ
                                1| : ||||: :| :
          309 IESYDLHFVKFAIDSDOLHOLWEKDOLNMAKNTWPILKGFQEGPLNFAPTFKFDVGTNKY 667
 כ'ת
          181 DTSEKKRKPAWTDRILWRLKRQPCAGPDTPIPPASH-FSLSLRGYSSHMTYGISDHKPVS 239
 C\lambda
              858 DTSAKKRKPAWTDRILWKVK-APGGGP-SPSGRKSHRLQVTQHSYRSHMEYTVSDHKPVA 725
· D5
          7.40 GTFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVGLRDVND 299
 Qу
                                         11: 1 1 1
          726 AQFLLQFAFRDDMPLVRLEVADEWVRPEQAVVRYRMETVFARSSWDWIGLYRVGFRHCKD 785
 Db
          300 YVSYAWVGDSKVSCSDNLNQVYIDISNIPTTEDEFLLCYYSNSLRSVVGISRPFQI 355
 Qу
                                             :1:1 | | | : :
                             | ||
                                     ::|
                                                        . : | | : | | | |
          786 YVAYVWAKHEDV--DGNTYQVTFSEESLPKGHGDFILGYYSHNHSILIGITEPFQI 839
 Db
 RESULT 8
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 DT
     01-MAR-2004 (TrEMBLrel. 26, Last sequence update)
 DT
     01-MAR-2004 (TrEMBLrel. 26, Last annotation update)
 DT
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     Name=acCG49651; ORFNames=ENSANGG00000016677;
 GΝ
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     Anopheles gambiae str. PEST.
     Eukaryota; Metazoa; Arthropoda; Hexapoda; Insecta; Pterygota;
 OC
     Neoptera; Endopterygota; Diptera; Nematocera; Culicoidea; Anopheles.
 OC
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     NCBI TaxID=180454;
 RN
      [1]
     SEQUENCE FROM N.A.
 RP
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STRAIN=PEST;
RA
    Anopheles Genome Sequencing Consortium;
    Submitted (MAR-2002) to the EMBL/GenBank/DDBJ databases.
RL
    -!- CAUTION: The sequence shown here is derived from an
CC
        EMBL/GenBank/DDBJ whole genome shotgun (WGS) entry which is
CC
CC
        preliminary data.
    EMBL; AAAB01008986; EAA00417.1; -.
DR
DR
    HSSP; 043001; 119Z.
    InterPro: IPR005135; Exo endo phos.
DR
    Pfam; PF03372; Exo_endo phos; 1.
DR
    NON TER
FT
    SEOUENCE
              501 AA; 57809 MW; F8C7DFBBF38FCA79 CRC64;
SO
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 Query Match
 Best Local Similarity 30.7%; Pred. No. 9e-36;
 Matches 116; Conservative 70; Mismatches 152; Indels
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                                                                   7;
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QУ
                 127 EVLKERDYVVIKTEQMQGLLLSVFARRKHLLHLRQVETEYTRTGLGGIWGNKGAVSIRMN 186
Db
         62 LYGYYVSIINCHLPPHISNNYQRLEHFDRILEMQNCEGRDIPNILDHDLIIWFGDMNFRI 121
Qу
            137 VYGSSICLVNAHLAAHDHMLEERINDYERIVQEQKFHVKAKETIFDHDYVFWFGDLNFRL 246
Db
       122 -----EDFGLHFVRESIKNRCYGGLWEKDQLSIAKKHDPLLREFQEGRLLFPPTYKF 173
Qу
                   : ||
        247 TGEATTSADEIRAMVARDELKQ-----LIEKDQLLLVRREGRAFAQLQERLPQFPPTFKF 301
Db
       174 DRNSNDYDTSEKKRKPAWTDRILWRLKRQPCAGPDTPIPPASHFSLSLRGYSSHMTYGIS 233 - ...
QΥ
                      ||:||||||||
            : ||:||
                                                  :
                                                        302 EHGSNEYD---MKRRPAWTDRILYAVNENNYRN------VRLTAEQTSYKSHPSYSIS 350
Db
        234 DHKPVSGTFDLELKPLVSAPLIVLMPEDLWTVENDMMVSYSSTSDFPSSPWDWIGLYKVG 293 -
Qу
            : | : |
        351 DHKPVTSEFTLKVYEDTTERTVSFKPIELWLVGEPNTIEYVIPNGFEEGNADWIGIYREN 410
Db
        294 LRDVNDYVSYAW------VGDSKVSCSDNLNQVYIDIS---NIPTTEDEFLLCYYS 340
Qу
               411 FTSLSEYLAYEYTETYKDKQHQLNEQQLQHMQNTRTVQLTFSENVSLPLGTRYQLLYFQS 470
Db
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    01-MAY-2000 (TrEMBLrel. 13, Last sequence update)
DT
    25-OCT-2004 (TrEMBLrel. 28, Last annotation update)
DT
    CG9784-PA (SD21513p).
DΕ
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OC
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RC

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OC
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OC
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     MEDLINE=20196006; PubMed=10731132; DOI=10.1126/science.287.5461.2185;
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RA
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     Borkova D., Botchan M.R., Bouck J., Brokstein P., Brottier P.,
     Burtis K.C., Busam D.A., Butler H., Cadieu E., Center A., Chandra I.,
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     Cherry J.M., Cawley S., Dahlke C., Davenport L.B., Davies P.,
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     de Pablos B., Delcher A., Deng Z., Mays A.D., Dew I., Dietz S.M.,
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     Dodson K., Doup L.E., Downes M., Dugan-Rocha S., Dunkov B.C., Dunn P.,
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     Durbin K.J., Evangelista C.C., Ferraz C., Ferriera S., Fleischmann W.,
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     Fosler C., Gabrielian A.E., Garg N.S., Gelbart W.M., Glasser K.,
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     Glodek A., Gong F., Gorrell J.H., Gu Z., Guan P., Harris M.,
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RR
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     Kimmel D.E., Kodira C.D., Kraft C., Kravitz S., Kulp D., Lai Z.,
     Basko P., Lei Y., Levitsky A.A., Li J., Li Z., Liang Y., Lin X.,
KK
     Liu K., Mattei B., McIntosh T.C., McLeod M.P., McPherson D.,
RA
     Merkulov G., Milshina N.V., Mobarry C., Morris J., Moshrefi A:,
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     Mount S.M., Moy M., Murphy B., Murphy L., Muzny D.M., Nelson D.L.,
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L\Omega
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RA
     Palazzolo M., Pittman G.S., Pan S., Pollard J., Puri V., Reese M.G.,
AS
     Reinert K., Remington K., Saunders R.D., Scheeler F., Shen H.,
     Shue B.C., Siden-Kiamos I., Simpson M., Skupski M.P., Smith T.,
RA
RΑ
     Spier E., Spradling A.C., Stapleton M., Strong R., Sun E.,
RA
     Svirskas R., Tector C., Turner R., Venter E., Wang A.H., Wang X.,
     Wang Z.Y., Wassarman D.A., Weinstock G.M., Weissenbach J.,
RA
     Williams S.M., WoodageT, Worley K.C., Wu D., Yang S., Yao Q.A., Ye J.,
RA
     Yeh R.F., Zaveri J.S., Zhan M., Zhang G., Zhao Q., Zheng L.,
RΆ
     Zheng X.H., Zhong F.N., Zhong W., Zhou X., Zhu S., Zhu X., Smith H.O.,
RA
     Gibbs R.A., Myers E.W., Rubin G.M., Venter J.C.;
RA
RT
     "The genome sequence of Drosophila melanogaster.";
RL
     Science 287:2185-2195(2000).
RN
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RX
     MEDLINE=22426065; PubMed=12537568;
     Celniker S.E., Wheeler D.A., Kronmiller B., Carlson J.W., Halpern A.,
RA
RA
     Patel S., Adams M., Champe M., Dugan S.P., Frise E., Hodgson A.,
RA
     George R.A., Hoskins R.A., Laverty T., Muzny D.M., Nelson C.R.,
     Pacleb J.M., Park S., Pfeiffer B.D., Richards S., Sodergren E.J.,
RΑ
     Svimskas R., Tabor P.E., Wan K., Stapleton M., Sutton G.G., Venter C.,
RA
     Weinstock G.. Scherer S.E., Myers E.W., Gibbs R.A., Rubin G.M.;
RA
     "Finishing a whole-genome shotgun: Release 3 of the Drosophila
RT
RT
     melanogaster euchromatic genome sequence.";
     Genome Biol. 3:RESEARCH0079-RESEARCH0079(2002).
RL
RN
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MEDLINE=22426070; PubMed=12537573;
RX
     Kaminker J.S., Bergman C.M., Kronmiller B., Carlson J., Svirskas R.,
RA
     Patel S., Frise E., Wheeler D.A., Lewis S.E., Rubin G.M.,
RA
     Ashburner M., Celniker S.E.;
RA
     "The transposable elements of the Drosophila melanogaster euchromatin:
RT
     a genomics perspective.";
RT
RL
     Genome Biol. 3:RESEARCH0084-RESEARCH0084(2002).
RN
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RA
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RA
     Smith C.D., Tupy J.L., Whitfied E.J., Bayraktaroglu L., Berman B.P.,
RA
RA
     Bettencourt B.R., Celniker S.E., de Grey A.D., Drysdale R.A.,
     Harris N.L., Richter J., Russo S., Schroeder A.J., Shu S.Q.,
RA
     Stapleton M., Yamada C., Ashburner M., Gelbart W.M., Rubin G.M.,
RA
RA
     "Annotation of the Drosophila melanogaster euchromatic genome: a
RT
RT
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RL
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RL
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RN
RР
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RG
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     submitted (MAR-2004) to the EMBI/GenBank/DDBJ databases.
RL
\mathbb{R}^{N}
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RΡ
L\Sigma
     Stapleton M., Brokstein P., Hong L., Agbayani A., Carlson J.,
\mathbb{R}^{p_{n}}
     Champe M., Chavez C., Dorsett V., Dresnek D., Farfan D., Frise E.,
RA. George R., Gonzalez M., Guarin H., Kronmiller B., Li P., Liao G.,
     Miranda A., Mungall C.J., Nunoo J., Pacleb J., Paragas V., Park S.,
RA
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     Submitted (JUN-2002) to the EMBL/GenBank/DDBJ databases.
RL
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     EMBL; AE003502; PAF48627.1; -.
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DR
     IntAct; Q9VXE7; -.
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     CO; GO:0004437; F:inositol or phosphatidylinositol phosphatas. . .; IEA.
DR
DR
     InterPro; IPR005135; Exo endo phos.
DR
     InterPro; IPR000300; IPPc.
DR
     Pfam; PF03372; Exo_endo_phos; 1.
DR
     SMART; SM00128; IPPc; 1.
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  Dost Local Similarity
                          34.3%;
                                  Pred. No. 5e-35;
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                                                                21;
                                                                     Gaps
                                                                             7;
            2 DVLSPLSFIKVSHVRMQGILLLVFAKYQHLPYIQILSTKSTPTGLFGYWGNKGGVNICLK 61
Qу
                           Db
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RΡ

SEQUENCE FROM N.A.

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62 LYGYYVSIINCHLPPHISNNYQRLEHFDRILEMQNCEGRDIPNILDHDLIIWFGDMNFRI 121
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          166 LYGCGLAFVVAHLTAHDHMMDERIEDYKQILENHHYHVKRYREIYDHDYVFWFGDLNFRL 225
Db
          122 E--DFGLHFVRESIKNRC-YGGLWEKDOL-SIAKKHDPLLREFOEGRLLFPPTYKFDRNS 177
Qу
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QУ
          295 RDVNDYVSYAWVGDSKVSCSDNLN 318
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Db
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DT
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     01-MAY-2000 (TrEMBLrel. 13, Last sequence update)
יבים
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ĐΤ
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DE
     CRFNames=CG6805;
GN
OS
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OC
     Bukaryota; Metazoa; Arthropoda; Hexapoda; Insecta; Pterygota;
OC
    Neoptera; Endopterygota; Diptera; Brachycera; Muscomorpha;
    Ephydroidea; Drosophilidae; Drosophila.
O.C
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OX
RN
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\mathbb{R}\mathbb{P}
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     MEDLINE=20196006; PubMed=10731132; DOI=10.1126/science.287.5461.2185;
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     Adams M.D., Celniker S.E., Holt R.A., Evans C.A., Gocayne J.D.,
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     Brandon R.C., Rogers Y.H., Blazej R.G., Champe M., Pfeiffer B.D.,
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     Wan K.H., Doyle C., Baxter E.G., Helt G., Nelson C.R., Gabor G.L.,
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ŔA
     Eceson K.Y., Benos P.V., Berman B.P., Bhandari D., Bolshakov S.,
     Borkova D., Botchan M.R., Bouck J., Brokstein P., Brottier P.,
RA
     Burtis K.C., Busam D.A., Butler H., Cadieu E., Center A., Chandra I.,
RA
     Cherry J.M., Cawley S., Dahlke C., Davenport L.B., Davies P.,
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    Jones S.J., Marra M.A.;
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"Generation and initial analysis of more than 15,000 full-length human
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    Jones S.J., Marra M.A.;
     "Generation and initial analysis of more than 15,000 full-length human
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     01-JUL-1993 (Rel. 26, Created)
DT
     01-FEB-1996 (Rel. 33, Last sequence update)
DT
DT
     05-JUL-2004 (Rel. 44, Last annotation update)
DE
     Type II inositol-1,4,5-trisphosphate 5-phosphatase precursor
DE
     (EC 3.1.3.36) (Phosphoinositide 5-phosphatase) (5PTase) (Fragment).
CN
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     Homo sapiens (Human).
OC
     Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
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RX
     NEDLINE=95238452; PubMed=7721860; DOI=10.1074/jbc.270.16.9370;
     Jefferson A.B., Majerus P.W.;
DA.
RT.
     "Properties of type II inositol polyphosphate 5-phosphatase.";
RL
     J. Biol. Chem. 270:9370-9377(1995).
RN
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     SEQUENCE OF 233-903 FROM N.A., AND SEQUENCE OF 270-288.
32
     TISSUE=Placenta;
RC
RX
     MEDLINE=92041857; PubMed=1718960;
     Ross T.S., Jefferson A.B., Mitchell C.A., Majerus P.W.;
RA
RT
    "Cloning and expression of human 75-kDa inositol polyphosphate-5-
RT
     phosphatase.";
     J. Biol. Chem. 266:20283-20289(1991).
P.L
     -!- FUNCTION: Hydrolyzes the calcium-mobilizing second messenger
CC
CC
         Ins(1,4,5)P3, this is a signal-terminating reaction.
CC
     -!- CATALYTIC ACTIVITY: 1-phosphatidyl-1D-myo-inositol 4,5-
CC
        bisphosphate + H(2)O = 1-phosphatidyl-1D-myo-inositol 4-phosphate
CC
         + phosphate.
CC --!- TISSUE SPECIFICITY: Platelets.
CC --!- SIMILARITY: Belongs to the inositol-1,4,5-trisphosphate 5-
CC .
        phosphatase type II family.
    -!- SIMILARITY: Contains 1 Rho-GAP domain.
CC
CC . -----
CC
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CC
     between the Swiss Institute of Bioinformatics and the EMBL outstation -
CC
     the European Bioinformatics Institute. There are no restrictions on its
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     entities requires a license agreement (See http://www.isb-sib.ch/announce/
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     or send an email to license@isb-sib.ch).
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EMBL; M74161; AAA79207.1; -.
DR
    HSSP; 043001; 119Z.
DR
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DR
DR
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    GO; GO:0004445; F:inositol-polyphosphate 5-phosphatase activity; NAS.
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DR
     InterPro; IPR000300; IPPc.
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DT
     01-MAR-2003 (TrEMBLrel. 23, Last sequence update)
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     01-OCT-2003 (TrEMBLrel. 25, Last annotation update)
     Mus musculus adult male lung cDNA, RIKEN full-length enriched library,
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     clone:1200004L23 product:inositol polyphosphate-5-phosphatase, 75 kDa,
DΕ
DΕ
     full insert sequence.
     Mus musculus (Mouse).
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RX
RA
     Carninci P., Hayashizaki Y.;
RT
     "High-efficiency full-length cDNA cloning.";
     Meth. Enzymol. 303:19-44(1999).
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     STRAIN=C57BL/6J; TISSUE=Lung;
     MEDLINE=21085660; PubMed=11217851; DOI=10.1038/35055500;
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RA
     RIKEN FANTOM Consortium;
RT
     "Functional annotation of a full-length mouse cDNA collection.";
RL
     Nature 409:685-690(2001).
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RC
     STRAIN=C57BL/6J; TISSUE=Lung;
RA
     The FANTOM Consortium,
RA
     the RIKEN Genome Exploration Research Group Phase I & II Team;
     "Analysis of the mouse transcriptome based on functional annotation of
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     Nature 420:563-573 (2002).
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     Carrinci P., Shibata Y., Hayatsu N., Sugahara Y., Shibata K., Itoh M.,
     Konno H., Okazaki Y., Muramatsu M., Hayashizaki Y.;
RA
RT
     "Normalization and subtraction of cap-trapper-selected cDNAs to
     prepare full-length cDNA libraries for rapid discovery of new genes.";
RT
RL
     Genome Res. 10:1617-1630(2000).
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     STRAIN=C57BL/6J; TISSUE=Lung;
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     Shibata K., Itoh M., Aizawa K., Nagaoka S., Sasaki N., Carninci P.,
RA
     Konno H., Akiyama J., Nishi K., Kitsunai T., Tashiro H., Itoh M.,
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     Sumi N., Ishii Y., Nakamura S., Hazama M., Nishine T., Harada A.,
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     Yamamoto R., Matsumoto H., Sakaguchi S., Ikegami T., Kashiwagi K.,
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     Fujiwake S., Inoue K., Togawa Y., Izawa M., Ohara E., Watahiki M.,
    'Yoneda Y., Ishikawa T., Ozawa K., Tanaka T., Matsuura S., Kawai J.,
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     Okazaki Y., Muramatsu M., Inoue Y., Kira A., Hayashizaki Y.;
     "RIKEN integrated sequence analysis (RISA) system-384-format
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     sequencing pipeline with 384 multicapillary sequencer.";
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     Genome Res. 10:1757-1771(2000).
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     Adachi J., Aizawa K., Akahira S., Akimura T., Arai A., Aono H.,
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Arakawa T., Bono H., Carninci P., Fukuda S., Fukunishi Y., Furuno M.,
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    Imotani K., Ishii Y., Itoh M., Izawa M., Kasukawa T., Kato H.,
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    Matsuyama T., Miyazaki A., Nishi K., Nomura K., Numazaki R., Ohno M.,
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    Okazaki Y., Okido T., Owa C., Saito H., Saito R., Sakai C., Sakai K.,
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    Sano H., Sasaki D., Shibata K., Shibata Y., Shinagawa A., Shiraki T.,
    Sogabe Y., Suzuki H., Tagami M., Tagawa A., Takahashi F., Tanaka T.,
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    Tejima Y., Toya T., Yamamura T., Yasunishi A., Yoshida K., Yoshino M.,
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    Muramatsu M., Hayashizaki Y.;
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    Submitted (JUL-2000) to the EMBL/GenBank/DDBJ databases.
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    InterPro; IPR000300; IPPc.
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    InterPro; IPR000198; RhoGAP.
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    InterPro; IPR008936; Rho GAP.
    Pfam; PF03372; Exo endo phos; 1.
DR
ÐR
    SMART; SM00128; IPPc; 1.
    SMART; SM00324; RhoGAP; 1.
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    PROSITE; PS50238; RHOGAP; 1.
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                                                                 Gaps
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Qy \cdot \cdot
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